

Lyme Disease Vaccines

5 Field of the Invention

The present invention relates to novel vaccines for the prevention or attenuation of Lyme disease. The invention further relates to isolated nucleic acid molecules encoding antigenic 10 polypeptides of *Borrelia burgdorferi*. Antigenic polypeptides are also provided, as are vectors, host cells and recombinant methods for producing the same. The invention additionally relates to diagnostic methods for detecting *Borrelia* gene expression.

15 45 Background of the Invention

Lyme disease (Steere, A.C., *Proc. Natl. Acad. Sci. USA* 91:2378-2383 (1991)), or Lyme borreliosis, is presently the most common human disease in the United States transmitted by an arthropod vector (Center for Disease Control, *Morbid. Mortal. Weekly Rep.* 46(23):531-535 (1997)). Further, infection of house-hold pets, such as dogs, is a considerable problem.

While initial symptoms often include a rash at the infection point, Lyme disease is a multisystemic disorder that may include arthritic, carditic, and neurological manifestations. While antibiotics are currently used to treat active cases of Lyme disease, *B. burgdorferi* persists even after prolonged antibiotic treatment. Further, *B. burgdorferi* can persist for years in a mammalian 25 host in the presence of an active immune response (Straubinger, R. *et al.*, *J. Clin. Microbiol.* 35:111-116 (1997); Steere, A., *N. Engl. J. Med.* 321:586-596 (1989)).

Lyme disease is caused by the related tick-borne spirochetes classified as *Borrelia burgdorferi* sensu lato (including *B. burgdorferi* sensu stricto, *B. afzelii*; *B. garinii*). Although substantial progress has been made in the biochemical, ultrastructural, and genetic characterization 30 of the organism, the spirochetal factors responsible for infectivity, immune evasion and disease pathogenesis remain largely obscure.

A number of antigenic *B. burgdorferi* cell surface proteins have been identified. These include the outer membrane surface proteins (Osp) OspA, OspB, OspC and OspD. OspA and OspB are encoded by tightly linked tandem genes which are transcribed as a single transcriptional 35 unit (Brusca, J. *et al.*, *J. Bacteriol.* 173:8004-8008 (1991)). The most-studied *B. burgdorferi* membrane protein is OspA, a lipoprotein antigen expressed by borreliae in resting ticks and the most abundant protein expressed *in vitro* by most borrelial isolates (Barbour, A.G., *et al.*, *Infection & Immunity* 41:795-804 (1983); Howe, T.R., *et al.*, *Science* 227:645 (1985)).

A number of different types of Lyme disease vaccines have been shown to induce immunological responses. Whole-cell *B. burgdorferi* vaccines, for example, have been shown to induce both immunological responses and protective immunity in several animal models.

5 (Reviewed in Wormser, G., *Clin. Infect. Dis.* 21:1267-1274 (1995)). Further, passive immunity has been demonstrated in both humans and other animals using *B. burgdorferi* specific antisera.

While whole-cell Lyme disease vaccines confer protective immunity in animal models, use of such vaccines presents the risk that responsive antibodies will produce an autoimmune response (Reviewed in Wormser, G., *supra*). This problem is at least partly the result of the production of *B. burgdorferi* specific antibodies which cross-react with hepatocytes and both 10 muscle and nerve cells. *B. burgdorferi* heat shock proteins and the 41-kd flagellin subunit are believed to contain antigens which elicit production of these cross-reactive antibodies.

Single protein subunit vaccines for Lyme disease have also been tested. The cell surface proteins of *B. burgdorferi* are potential candidates for use in such vaccines and several have been shown to elicit protective immune responses in mammals (Probert, W., *et al.*, *Vaccine* 15:15-19 (1997); Fikrig, E., *et al.*, *Infect. Immun.* 63:1658-1662 (1995); Langerman S., *et al.*, *Nature* 372:552-556 (1994); Fikrig, E., *et al.*, *J. Immunol.* 148:2256-2260 (1992)). Experimental OspA vaccines, for example, have demonstrated efficacy in several animal models (Fikrig, E., *et al.*, *Proc. Natl. Acad. Sci. USA* 89:5418-5421 (1992); Johnson, B.J., *et al.*, *Vaccine* 13:1086-1094 (1996); Fikrig, E., *et al.*, *Infect. Immun.* 60:657-661 (1992); Chang, Y.F., *et al.*, *Infection & Immunity* 63:3543-3549 (1995)), and OspA vaccines for human use are under clinical evaluation (Keller, D., *et al.*, *J. Am. Med. Assoc.* 271:1764-1768 (1994); Van Hoecke, C., *et al.*, *Vaccine* 14:1620-1626 (1996)). Passive immunity is also conferred by antisera containing antibodies specific for the full-length OspA protein. Further, vaccination with plasmid DNA encoding OspA has been demonstrated to elicit protective immune responses in mice (Luke, C., *et al.*, *J. Infect. Dis.* 175:91-97 (1997); Zhong, W., *et al.*, *Eur. J. Immunol.* 26:2749-2757 (1996)).

Recent immunofluorescence assay observations indicate that during tick engorgement the expression of OspA by borreliae diminishes (deSilva, A.M., *et al.*, *J. Exp. Med.* 183:271-275 (1996)) while expression of other proteins, exemplified by OspC, increases (Schwan, T.G., *et al.*, *Proc. Natl. Acad. Sci. USA* 92:2909-2913 (1985)). By the time of transmission to hosts, 30 spirochetes in the tick salivary glands express little or no OspA. This down-modulation of OspA appears to explain the difficulties in demonstrating immune responses to this antigen early in infection following tick bites (Kalish, R.A., *et al.*, *Infect. Immun.* 63:2228-2235 (1995); Gern, L., *et al.*, *J. Infect. Dis.* 167:971-975 (1993); Schiable, U.E., *et al.*, *Immunol. Lett.* 36:219-226 (1993)) or following challenge with limiting doses of cultured borreliae (Schiable, U.E., *et al.*, *Immunol. Lett.* 36:219-226 (1993); Barthold, S.W. and Bockenstedt, L.K., *Infect. Immun.* 61:4696-4702 (1993)).

Furthermore, OspA-specific antibodies are ineffective if administered after a borrelial challenge delivered by syringe (Schiable, U.E., *et al.*, *Proc. Natl. Acad. Sci. USA* 87:3768-3772 (1990)) or tick bite (deSilva, A.M., *et al.*, *J. Exp. Med.* 183:271-275 (1996)). To be efficacious,

OspA vaccines must elicit protective levels of antibody which are maintained throughout periods of tick exposure in order to block borrelia transmission from the arthropod vector.

Vaccines in current use against other pathogens include *in vivo*-expressed antigens which could boost anamnestic responses upon infection, potentiate the action of immune effector cells and complement, and inhibit key virulence mechanisms. OspC is both expressed during infection (Montgomery, R.R., *et al.*, *J. Exp. Med.* 183:261-269 (1996)) and a target for protective immunity (Gilmore, R.D., *et al.*, *Infect. Immun.* 64:2234-2239 (1996); Probert, W.S. and LeFebvre, R.B., *Infect. Immun.* 62:1920-1926 (1994); Preac-Mursic, V., *et al.*, *Infection* 20:342-349 (1992)), but mice immunized with this protein were only protected against challenge with the homologous borrelial isolate (Probert, W.S., *et al.*, *J. Infect. Dis.* 175:400-405 (1997)). Identification of *in vivo*-expressed, and broadly protective, antigens of *B. burgdorferi* has remained elusive.

Summary of the Invention

The present invention provides isolated nucleic acid molecules comprising polynucleotides encoding the *B. burgdorferi* peptides having the amino acid sequences shown in Table 1. Thus, one aspect of the invention provides isolated nucleic acid molecules comprising polynucleotides having a nucleotide sequence selected from the group consisting of: (a) a nucleotide sequence encoding any of the amino acid sequences of the full-length polypeptides shown in Table 1; (b) a nucleotide sequence encoding any of the amino acid sequences of the full-length polypeptides shown in Table 1 but minus the N-terminal methionine residue, if present; (c) a nucleotide sequence encoding any of the amino acid sequences of the truncated polypeptides shown in Table 1; and (d) a nucleotide sequence complementary to any of the nucleotide sequences in (a), (b), or (c) above.

Further embodiments of the invention include isolated nucleic acid molecules that comprise a polynucleotide having a nucleotide sequence at least 90% identical, and more preferably at least 95%, 96%, 97%, 98% or 99% identical, to any of the nucleotide sequences in (a), (b), (c), or (d) above, or a polynucleotide which hybridizes under stringent hybridization conditions to a polynucleotide in (a), (b), (c), or (d) above. This polynucleotide which hybridizes does not hybridize under stringent hybridization conditions to a polynucleotide having a nucleotide sequence consisting of only A residues or of only T residues. Additional nucleic acid embodiments of the invention relate to isolated nucleic acid molecules comprising polynucleotides which encode the amino acid sequences of epitope-bearing portions of a *B. burgdorferi* polypeptide having an amino acid sequence in (a), (b), or (c) above.

The present invention also relates to recombinant vectors, which include the isolated nucleic acid molecules of the present invention, and to host cells containing the recombinant vectors, as well as to methods of making such vectors and host cells and for using these vectors for the production of *B. burgdorferi* polypeptides or peptides by recombinant techniques.

The invention further provides isolated *B. burgdorferi* polypeptides having an amino acid

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sequence selected from the group consisting of: (a) an amino acid sequence of any of the full-length polypeptides shown in Table 1; (b) an amino acid sequence of any of the full-length polypeptides shown in Table 1 but minus the N-terminal methionine residue, if present; (c) an amino acid sequence of any of the truncated polypeptides shown in Table 1; and (d) an amino acid sequence of an epitope-bearing portion of any one of the polypeptides of (a), (b), or (c).

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The polypeptides of the present invention also include polypeptides having an amino acid sequence with at least 70% similarity, and more preferably at least 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98%, or 99% similarity to those described in (a), (b), (c), or (d) above, as well as polypeptides having an amino acid sequence at least 70% identical, more preferably at least 75% identical, and still more preferably 80%, 85%, 90%, 95%, 96%, 97%, 98%, or 99% identical to those above; as well as isolated nucleic acid molecules encoding such polypeptides.

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The present invention further provides a vaccine, preferably a multi-component vaccine comprising one or more of the *B. burgdorferi* polypeptides shown in Table 1, or fragments thereof, together with a pharmaceutically acceptable diluent, carrier, or excipient, wherein the *B. burgdorferi* polypeptide(s) are present in an amount effective to elicit an immune response to members of the *Borrelia* genus in an animal. The *B. burgdorferi* polypeptides of the present invention may further be combined with one or more immunogens of one or more other borrelial or non-borrelial organisms to produce a multi-component vaccine intended to elicit an immunological response against members of the *Borrelia* genus and, optionally, one or more non-borrelial organisms.

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The vaccines of the present invention can be administered in a DNA form, e.g., "naked" DNA, wherein the DNA encodes one or more borrelial polypeptides and, optionally, one or more polypeptides of a non-borrelial organism. The DNA encoding one or more polypeptides may be constructed such that these polypeptides are expressed fusion proteins.

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The vaccines of the present invention may also be administered as a component of a genetically engineered organism. Thus, a genetically engineered organism which expresses one or more *B. burgdorferi* polypeptides may be administered to an animal. For example, such a genetically engineered organism may contain one or more *B. burgdorferi* polypeptides of the present invention intracellularly, on its cell surface, or in its periplasmic space. Further, such a genetically engineered organism may secrete one or more *B. burgdorferi* polypeptides.

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The vaccines of the present invention may be co-administered to an animal with an immune system modulator (e.g., CD86 and GM-CSF).

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The invention also provides a method of inducing an immunological response in an animal to one or more members of the *Borrelia* genus, e.g., *B. burgdorferi* sensu stricto, *B. afzelii*, and *B. garinii*, comprising administering to the animal a vaccine as described above.

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The invention further provides a method of inducing a protective immune response in an animal, sufficient to prevent or attenuate an infection by members of the *Borrelia* genus, comprising administering to the animal a composition comprising one or more of the polypeptides shown in Table 1, or fragments thereof. Further, these polypeptides, or fragments thereof, may

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be conjugated to another immunogen and/or administered in admixture with an adjuvant.

The invention further relates to antibodies elicited in an animal by the administration of one or more *B. burgdorferi* polypeptides of the present invention.

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The invention also provides diagnostic methods for detecting the expression of genes of members of the *Borrelia* genus in an animal. One such method involves assaying for the expression of a gene encoding *Borrelia* peptides in a sample from an animal. This expression may be assayed either directly (e.g., by assaying polypeptide levels using antibodies elicited in response to amino acid sequences shown in Table 1) or indirectly (e.g., by assaying for antibodies having specificity for amino acid sequences shown in Table 1). An example of such a method involves the use of the polymerase chain reaction (PCR) to amplify and detect *Borrelia* nucleic acid sequences.

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The present invention also relates to nucleic acid probes having all or part of a nucleotide sequence shown in Table 1 which are capable of hybridizing under stringent conditions to *Borrelia* nucleic acids. The invention further relates to a method of detecting one or more *Borrelia* nucleic acids in a biological sample obtained from an animal, said one or more nucleic acids encoding *Borrelia* polypeptides, comprising:

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a) contacting the sample with one or more of the above-described nucleic acid probes, under conditions such that hybridization occurs, and
b) detecting hybridization of said one or more probes to the *Borrelia* nucleic acid present in the biological sample.

25 **Detailed Description**

The present invention relates to recombinant antigenic *B. burgdorferi* polypeptides and fragments thereof. The invention also relates to methods for using these polypeptides to produce immunological responses and to confer immunological protection to disease caused by members 30
of the genus *Borrelia*. The invention further relates to nucleic acid sequences which encode antigenic *B. burgdorferi* polypeptides and to methods for detecting *Borrelia* nucleic acids and polypeptides in biological samples. The invention also relates to *Borrelia* specific antibodies and methods for detecting such antibodies produced in a host animal.

35 **Definitions**

The following definitions are provided to clarify the subject matter which the inventors consider to be the present invention.

As used herein, the phrase "pathogenic agent" means an agent which causes a disease state or affliction in an animal. Included within this definition, for examples, are bacteria, protozoans, fungi, viruses and metazoan parasites which either produce a disease state or render an animal infected with such an organism susceptible to a disease state (e.g., a secondary infection). Further included are species and strains of the genus *Borrelia* which produce disease states in animals.

As used herein, the term "organism" means any living biological system, including viruses, regardless of whether it is a pathogenic agent.

As used herein, the term "Borrelia" means any species or strain of bacteria which is members of the genus *Borrelia*. Included within this definition are *Borrelia burgdorferi* sensu lato (including *B. burgdorferi* sensu stricto, *B. afzelii*, *B. garinii*), *B. andersonii*, *B. anserina*, *B. japonica*, *B. coriaceae*, and other members of the genus *Borrelia* regardless of whether they are known pathogenic agents.

As used herein, the phrase "one or more *B. burgdorferi* polypeptides of the present invention" means the amino acid sequence of one or more of the *B. burgdorferi* polypeptides disclosed in Table 1. These polypeptides may be expressed as fusion proteins wherein the *B. burgdorferi* polypeptides of the present invention are linked to additional amino acid sequences which may be of borrelial or non-borrelial origin. This phrase further includes fragments of the *B. burgdorferi* polypeptides of the present invention.

As used herein, the phrase "full-length amino acid sequence" and "full-length polypeptide" refer to an amino acid sequence or polypeptide encoded by a full-length open reading frame (ORF). An ORF may be defined as a nucleotide sequence bounded by stop codons which encodes a putative polypeptide. An ORF may also be defined as a nucleotide sequence within a stop codon bounded sequence which contains an initiation codon (e.g., a methionine or valine codon) on the 5' end and a stop codon on the 3' end.

As used herein, the phrase "truncated amino acid sequence" and "truncated polypeptide" refer to a sub-sequence of a full-length amino acid sequence or polypeptide. Several criteria may also be used to define the truncated amino acid sequence or polypeptide. For example, a truncated polypeptide may be defined as a mature polypeptide (e.g., a polypeptide which lacks a leader sequence). A truncated polypeptide may also be defined as an amino acid sequence which is a portion of a longer sequence that has been selected for ease of expression in a heterologous system but retains regions which render the polypeptide useful for use in vaccines (e.g., antigenic regions which are expected to elicit a protective immune response).

Additional definitions are provided throughout the specification.

Explanation of Table 1

Table 1 lists *B. burgdorferi* nucleotide and amino acid sequences of the present invention. The nomenclature used therein is as follows:

"nt" refers to nucleotide sequences;

"aa" refers to amino acid sequences;

"f" refers to full-length nucleotide or amino acid sequences; and

"t" refers to truncated nucleotide or amino acid sequences.

Thus, for example, the designation "f101.aa" refers to the full-length amino acid sequence of *B. burgdorferi* polypeptide number 101. Further, "f101.nt" refers to the full-length nucleotide sequence encoding the full-length amino acid sequence of *B. burgdorferi* polypeptide number 101.

Explanation of Table 2

Table 2 lists accession numbers for the closest matching sequences between the polypeptides of the present invention and those available through GenBank and GeneSeq databases. These reference numbers are the database entry numbers commonly used by those of skill in the art, who will be familiar with their denominations. The descriptions of the nomenclature for GenBank are available from the National Center for Biotechnology Information. Column 1 lists the gene or ORF of the present invention. Column 2 lists the accession number of a "match" gene sequence in GenBank or GeneSeq databases. Column 3 lists the description of the "match" gene sequence. Columns 4 and 5 are the high score and smallest sum probability, respectively, calculated by BLAST. Polypeptides of the present invention that do not share significant identity/similarity with any polypeptide sequences of GenBank and GeneSeq are not represented in Table 2. Polypeptides of the present invention that share significant identity/similarity with more than one of the polypeptides of GenBank and GeneSeq are represented more than once.

Explanation of Table 3.

The *B. burgdorferi* polypeptides of the present invention may include one or more conservative amino acid substitutions from natural mutations or human manipulation as indicated in Table 3. Changes are preferably of a minor nature, such as conservative amino acid substitutions that do not significantly affect the folding or activity of the protein. Residues from the following groups, as indicated in Table 3, may be substituted for one another: Aromatic, Hydrophobic, Polar, Basic, Acidic, and Small.

Explanation of Table 4

Table 4 lists residues comprising antigenic epitopes of antigenic epitope-bearing fragments present in each of the full length *B. burgdorferi* polypeptides described in Table 1 as predicted by the inventors using the algorithm of Jameson and Wolf, (1988) Comp. Appl. Biosci. 4:181-186. The Jameson-Wolf antigenic analysis was performed using the computer program PROTEAN (Version 3.11 for the Power MacIntosh, DNASTAR, Inc., 1228 South Park Street Madison, WI). *B. burgdorferi* polypeptide shown in Table 1 may one or more antigenic epitopes comprising residues described in Table 4. It will be appreciated that depending on the analytical criteria used to predict antigenic determinants, the exact address of the determinant may vary slightly. The residues and locations shown described in Table 4 correspond to the amino acid sequences for each full length gene sequence shown in Table 1 and in the Sequence Listing. Polypeptides of the present invention that do not have antigenic epitopes recognized by the Jameson-Wolf algorithm are not represented in Table 2.

*Selection of Nucleic Acid Sequences Encoding Antigenic *B. burgdorferi* Polypeptides*

The present invention provides a select number of ORFs from those presented in the fragments of the *Borrelia burgdorferi* genome which may prove useful for the generation of a protective immune response. The sequenced *B. burgdorferi* genomic DNA was obtained from a sub-cultured isolate of ATCC Deposit No. 35210. The sub-cultured isolate was deposited on August 8, 1997 at the American Type Culture Collection, 12301 Park Lawn Drive, Rockville, Maryland 20852, and given accession number 202012.

Some ORFs contained in the subset of fragments of the *B. burgdorferi* genome disclosed herein were derived through the use of a number of screening criteria detailed below. The ORFs are generally bounded at the amino terminus by a methionine residue and at the carboxy terminus by a stop codon.

Many of the selected sequences do not consist of complete ORFs. Although a polypeptide representing a complete ORF may be the closest approximation of a protein native to an organism, it is not always preferred to express a complete ORF in a heterologous system. It may be challenging to express and purify a highly hydrophobic protein by common laboratory methods. Some of the polypeptide vaccine candidates described herein have been modified slightly to simplify the production of recombinant protein. For example, nucleotide sequences which encode highly hydrophobic domains, such as those found at the amino terminal signal sequence, have been excluded from some constructs used for *in vitro* expression of the polypeptides. Furthermore, any highly hydrophobic amino acid sequences occurring at the carboxy terminus have also been excluded from the recombinant expression constructs. Thus, in one embodiment, a polypeptide which represents a truncated or modified ORF may be used as an antigen.

While numerous methods are known in the art for selecting potentially immunogenic polypeptides, many of the ORFs disclosed herein were selected on the basis of screening all theoretical *Borrelia burgdorferi* ORFs for several aspects of potential immunogenicity. One set of selection criteria are as follows:

1. *Type I signal sequence*: An amino terminal type I signal sequence generally directs a nascent protein across the plasma and outer membranes to the exterior of the bacterial cell. Experimental evidence obtained from studies with *Escherichia coli* suggests that the typical type I signal sequence consists of the following biochemical and physical attributes (Izard, J. W. and Kendall, D. A. *Mol. Microbiol.* 13:765-773 (1994)). The length of the type I signal sequence is approximately 15 to 25 primarily hydrophobic amino acid residues with a net positive charge in the extreme amino terminus. In addition, the central region of the signal sequence adopts an alpha-helical conformation in a hydrophobic environment. Finally, the region surrounding the actual site of cleavage is ideally six residues long, with small side-chain amino acids in the -1 and -3 positions.

2. *Type IV signal sequence*: The type IV signal sequence is an example of the several types of functional signal sequences which exist in addition to the type I signal sequence detailed

above. Although functionally related, the type IV signal sequence possesses a unique set of biochemical and physical attributes (Strom, M. S. and Lory, S., *J. Bacteriol.* 174:7345-7351 (1992)). These are typically six to eight amino acids with a net basic charge followed by an additional sixteen to thirty primarily hydrophobic residues. The cleavage site of a type IV signal sequence is typically after the initial six to eight amino acids at the extreme amino terminus. In addition, type IV signal sequences generally contain a phenylalanine residue at the +1 site relative to the cleavage site.

3. *Lipoprotein*: Studies of the cleavage sites of twenty-six bacterial lipoprotein precursors has allowed the definition of a consensus amino acid sequence for lipoprotein cleavage. Nearly 10 three-fourths of the bacterial lipoprotein precursors examined contained the sequence L-(A,S)-(G,A)-C at positions -3 to +1, relative to the point of cleavage (Hayashi, S. and Wu, H. C., *J. Bioenerg. Biomembr.* 22:451-471 (1990)).

4. *LPXTG motif*: It has been experimentally determined that most anchored proteins found on the surface of gram-positive bacteria possess a highly conserved carboxy terminal sequence. More than fifty such proteins from organisms such as *S. pyogenes*, *S. mutans*, *B. burgdorferi*, *S. pneumoniae*, and others, have been identified based on their extracellular location and carboxy terminal amino acid sequence (Fischetti, V. A., *ASM News* 62:405-410 (1996)). The conserved region consists of six charged amino acids at the extreme carboxy terminus coupled to 15-20 hydrophobic amino acids presumed to function as a transmembrane domain. Immediately adjacent to the transmembrane domain is a six amino acid sequence conserved in nearly all proteins examined. The amino acid sequence of this region is L-P-X-T-G-X, where X is any amino acid.

An algorithm for selecting antigenic and immunogenic *Borrelia burgdorferi* polypeptides including the foregoing criteria was developed. The algorithm is similar to that described in U.S. 25 patent application 08/781,986, filed January 3, 1997, which is fully incorporated by reference herein. Use of the algorithm by the inventors to select immunologically useful *Borrelia burgdorferi* polypeptides resulted in the selection of a number of the disclosed ORFs. Polypeptides comprising the polypeptides identified in this group may be produced by techniques standard in the art and as further described herein.

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Nucleic Acid Molecules

The present invention provides isolated nucleic acid molecules comprising polynucleotides encoding the *B. burgdorferi* polypeptides having the amino acid sequences shown in Table 1, which were determined by sequencing the genome of *B. burgdorferi* deposited as ATCC deposit 35 no. 202012 and selected as putative immunogens.

Unless otherwise indicated, all nucleotide sequences determined by sequencing a DNA molecule herein were determined using an automated DNA sequencer (such as the Model 373 from Applied Biosystems, Inc.), and all amino acid sequences of polypeptides encoded by DNA molecules determined herein were predicted by translation of DNA sequences determined as

10 above. Therefore, as is known in the art for any DNA sequence determined by this automated approach, any nucleotide sequence determined herein may contain some errors. Nucleotide sequences determined by automation are typically at least about 90% identical, more typically at least about 95% to at least about 99.9% identical to the actual nucleotide sequence of the sequenced DNA molecule. The actual sequence can be more precisely determined by other approaches including manual DNA sequencing methods well known in the art. As is also known in the art, a single insertion or deletion in a determined nucleotide sequence compared to the actual sequence will cause a frame shift in translation of the nucleotide sequence such that the predicted amino acid sequence encoded by a determined nucleotide sequence will be completely different from the amino acid sequence actually encoded by the sequenced DNA molecule, beginning at the point of such an insertion or deletion.

Unless otherwise indicated, each "nucleotide sequence" set forth herein is presented as a sequence of deoxyribonucleotides (abbreviated A, G, C and T). However, by "nucleotide sequence" of a nucleic acid molecule or polynucleotide is intended, for a DNA molecule or polynucleotide, a sequence of deoxyribonucleotides, and for an RNA molecule or polynucleotide, the corresponding sequence of ribonucleotides (A, G, C and U), where each thymidine deoxyribonucleotide (T) in the specified deoxyribonucleotide sequence is replaced by the ribonucleotide uridine (U). For instance, reference to an RNA molecule having a sequence of Table 1 set forth using deoxyribonucleotide abbreviations is intended to indicate an RNA molecule having a sequence in which each deoxyribonucleotide A, G or C of Table 1 has been replaced by the corresponding ribonucleotide A, G or C, and each deoxyribonucleotide T has been replaced by a ribonucleotide U.

Nucleic acid molecules of the present invention may be in the form of RNA, such as mRNA, or in the form of DNA, including, for instance, cDNA and genomic DNA obtained by cloning or produced synthetically. The DNA may be double-stranded or single-stranded. Single-stranded DNA or RNA may be the coding strand, also known as the sense strand, or it may be the non-coding strand, also referred to as the anti-sense strand.

By "isolated" nucleic acid molecule(s) is intended a nucleic acid molecule, DNA or RNA, which has been removed from its native environment. For example, recombinant DNA molecules contained in a vector are considered isolated for the purposes of the present invention. Further examples of isolated DNA molecules include recombinant DNA molecules maintained in heterologous host cells or purified (partially or substantially) DNA molecules in solution. Isolated RNA molecules include *in vivo* or *in vitro* RNA transcripts of the DNA molecules of the present invention. Isolated nucleic acid molecules according to the present invention further include such molecules produced synthetically.

In addition, isolated nucleic acid molecules of the invention include DNA molecules which comprise a sequence substantially different from those described above but which, due to the degeneracy of the genetic code, still encode a *B. burgdorferi* polypeptides and peptides of the present invention (e.g. polypeptides of Table 1). That is, all possible DNA sequences that encode

the *B. burgdorferi* polypeptides of the present invention. This includes the genetic code and species-specific codon preferences known in the art. Thus, it would be routine for one skilled in the art to generate the degenerate variants described above, for instance, to optimize codon expression for a particular host (e.g., change codons in the bacteria mRNA to those preferred by a mammalian or other bacterial host such as *E. coli*).

The invention further provides isolated nucleic acid molecules having the nucleotide sequence shown in Table 1 or a nucleic acid molecule having a sequence complementary to one of the above sequences. Such isolated molecules, particularly DNA molecules, are useful as probes for gene mapping and for identifying *B. burgdorferi* in a biological sample, for instance, by PCR, Southern blot, Northern blot, or other form of hybridization analysis.

The present invention is further directed to nucleic acid molecules encoding portions or fragments of the nucleotide sequences described herein. Fragments include portions of the nucleotide sequences of Table 1 at least 10 contiguous nucleotides in length selected from any two integers, one of which representing a 5' nucleotide position and a second of which representing a 3' nucleotide position, where the first nucleotide for each nucleotide sequence in Table 1 is position 1. That is, every combination of a 5' and 3' nucleotide position that a fragment at least 10 contiguous nucleotides in length could occupy is included in the invention. "At least" means a fragment may be 10 contiguous nucleotide bases in length or any integer between 10 and the length of an entire nucleotide sequence of Table 1 minus 1. Therefore, included in the invention are contiguous fragments specified by any 5' and 3' nucleotide base positions of a nucleotide sequences of Table 1 wherein the contiguous fragment is any integer between 10 and the length of an entire nucleotide sequence minus 1.

Further, the invention includes polynucleotides comprising fragments specified by size, in nucleotides, rather than by nucleotide positions. The invention includes any fragment size, in contiguous nucleotides, selected from integers between 10 and the length of an entire nucleotide sequence minus 1. Preferred sizes of contiguous nucleotide fragments include 20 nucleotides, 30 nucleotides, 40 nucleotides, 50 nucleotides. Other preferred sizes of contiguous nucleotide fragments, which may be useful as diagnostic probes and primers, include fragments 50-300 nucleotides in length which include, as discussed above, fragment sizes representing each integer between 50-300. Larger fragments are also useful according to the present invention corresponding to most, if not all, of the nucleotide sequences shown in Table 1 or of the *B. burgdorferi* nucleotide sequences of the plasmid clones listed in Table 1. The preferred sizes are, of course, meant to exemplify not limit the present invention as all size fragments, representing any integer between 10 and the length of an entire nucleotide sequence minus 1, are included in the invention. Additional preferred nucleic acid fragments of the present invention include nucleic acid molecules encoding epitope-bearing portions of *B. burgdorferi* polypeptides identified in Table 4.

The present invention also provides for the exclusion of any fragment, specified by 5' and 3' base positions or by size in nucleotide bases as described above for any nucleotide sequence of

12 Table 1 or the plasmid clones listed in Table 1. Any number of fragments of nucleotide sequences in Table 1 or the plasmid clones listed in Table 1, specified by 5' and 3' base positions or by size in nucleotides, as described above, may be excluded from the present invention.

5 Preferred nucleic acid fragments of the present invention also include nucleic acid molecules encoding epitope-bearing portions of the *B. burgdorferi* polypeptides shown in Table 1. Such nucleic acid fragments of the present invention include, for example, nucleic acid molecules encoding polypeptide fragments comprising from about the amino terminal residue to about the carboxy terminal residue of each fragment shown in Table 4. The above referred to polypeptide fragments are antigenic regions of particular *B. burgdorferi* polypeptides shown in Table 1. Methods for determining other such epitope-bearing portions for the remaining polypeptides described in Table 1 are well known in the art and are described in detail below.

10 15 20 25 30 35 40 45 50 55 60 65 70 75 80 85 90 95 In another aspect, the invention provides isolated nucleic acid molecules comprising polynucleotides which hybridize under stringent hybridization conditions to a portion of a polynucleotide in a nucleic acid molecule of the invention described above, for instance, a nucleic acid sequence shown in Table 1. By "stringent hybridization conditions" is intended overnight incubation at 42 C in a solution comprising: 50% formamide, 5x SSC (150 mM NaCl, 15 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5x Denhardt's solution, 10% dextran sulfate, and 20 g/ml denatured, sheared salmon sperm DNA, followed by washing the filters in 0.1x SSC at about 65 C.

20 25 30 35 40 45 50 55 60 65 70 75 80 85 90 95 By polynucleotides which hybridize to a "portion" of a polynucleotide is intended polynucleotides (either DNA or RNA) which hybridize to at least about 15 nucleotides (nt), and more preferably at least about 20 nt, still more preferably at least about 30 nt, and even more preferably about 30-70 nt of the reference polynucleotide. These are useful as diagnostic probes and primers as discussed above and in more detail below.

25 30 35 40 45 50 55 60 65 70 75 80 85 90 95 Of course, polynucleotides hybridizing to a larger portion of the reference polynucleotide, for instance, a portion 50-100 nt in length, or even to the entire length of the reference polynucleotide, are also useful as probes according to the present invention, as are polynucleotides corresponding to most, if not all, of a nucleotide sequence as shown in Table 1. By a portion of a polynucleotide of "at least 20 nt in length," for example, is intended 20 or more contiguous nucleotides from the nucleotide sequence of the reference polynucleotide (e.g., a nucleotide sequence as shown in Table 1). As noted above, such portions are useful diagnostically either as probes according to conventional DNA hybridization techniques or as primers for amplification of a target sequence by PCR, as described, for instance, in *Molecular Cloning, A Laboratory Manual*, 2nd. edition, Sambrook, J., Fritsch, E. F. and Maniatis, T., eds., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (1989), the entire disclosure of which is hereby incorporated herein by reference.

30 35 40 45 50 55 60 65 70 75 80 85 90 95 Since nucleic acid sequences encoding the *B. burgdorferi* polypeptides of the present invention are provided in Table 1, generating polynucleotides which hybridize to portions of these sequences would be routine to the skilled artisan. For example, the hybridizing polynucleotides

of the present invention could be generated synthetically according to known techniques.

As indicated, nucleic acid molecules of the present invention which encode *B. burgdorferi* polypeptides of the present invention may include, but are not limited to those encoding the amino acid sequences of the polypeptides by themselves; and additional coding sequences which code for additional amino acids, such as those which provide additional functionalities. Thus, the sequences encoding these polypeptides may be fused to a marker sequence, such as a sequence encoding a peptide which facilitates purification of the fused polypeptide. In certain preferred embodiments of this aspect of the invention, the marker amino acid sequence is a hexa-histidine peptide, such as the tag provided in a pQE vector (Qiagen, Inc.), among others, many of which are commercially available. As described in Gentz *et al.*, *Proc. Natl. Acad. Sci. USA* 86:821-824 (1989), for instance, hexa-histidine provides for convenient purification of the resulting fusion protein.

Thus, the present invention also includes genetic fusions wherein the *B. burgdorferi* nucleic acid sequences coding sequences provided in Table 1 are linked to additional nucleic acid sequences to produce fusion proteins. These fusion proteins may include epitopes of borrelial or non-borrelial origin designed to produce proteins having enhanced immunogenicity. Further, the fusion proteins of the present invention may contain antigenic determinants known to provide helper T-cell stimulation, peptides encoding sites for post-translational modifications which enhance immunogenicity (e.g., acylation), peptides which facilitate purification (e.g., histidine "tag"), or amino acid sequences which target the fusion protein to a desired location (e.g., a heterologous leader sequence). For instance, hexa-histidine provides for convenient purification of the fusion protein. See Gentz *et al.* (1989) *Proc. Natl. Acad. Sci.* 86:821-24. The "HA" tag is another peptide useful for purification which corresponds to an epitope derived from the influenza hemagglutinin protein. See Wilson *et al.* (1984) *Cell* 37:767. As discussed below, other such fusion proteins include the *B. burgdorferi* polypeptides of the present invention fused to Fc at the N- or C-terminus.

Post-translational modification of the full-length *B. burgdorferi* OspA protein expressed in *E. coli* is believed to increase the immunogenicity of this protein. Erdile, L. *et al.*, *Infect. Immun.* 61:81-90 (1993). *B. burgdorferi* OspA when expressed in *E. coli*, for example, is post-translationally modified in at least two ways. First, a signal peptide is cleaved; second, lipid moieties are attached. The presence of these lipid moieties is believed to confer enhanced immunogenicity and results in the elicitation of a strong protective immunological response.

Variant and Mutant Polynucleotides

The present invention thus includes nucleic acid molecules and sequences which encode fusion proteins comprising one or more *B. burgdorferi* polypeptides of the present invention fused to an amino acid sequence which allows for post-translational modification to enhance immunogenicity. This post-translational modification may occur either *in vitro* or when the fusion protein is expressed *in vivo* in a host cell. An example of such a modification is the introduction

of an amino acid sequence which results in the attachment of a lipid moiety. Such a lipid moiety attachment site of OspA, which is lipidated upon expression in *E. coli*, has been identified. Bouchon, B. et al., *Anal. Biochem.* 246:52-61 (1997).

Thus, as indicated above, the present invention includes genetic fusions wherein a *B. burgdorferi* nucleic acid sequence provided in Table 1 is linked to a nucleotide sequence encoding another amino acid sequence. These other amino acid sequences may be of borrelial origin (e.g., another sequence selected from Table 1) or non-borrelial origin. An example of such a fusion protein is reported in Fikrig, E. et al., *Science* 250:553-556 (1990) where an OspA-glutathione-S-transferase fusion protein was produced and shown to elicit protective immunity against Lyme disease in immune competent mice.

The present invention further relates to variants of the nucleic acid molecules of the present invention, which encode portions, analogs or derivatives of the *B. burgdorferi* polypeptides shown in Table 1. Variants may occur naturally, such as a natural allelic variant. By an "allelic variant" is intended one of several alternate forms of a gene occupying a given locus on a chromosome of an organism. *Genes II*, Lewin, B., ed., John Wiley & Sons, New York (1985). Non-naturally occurring variants may be produced using art-known mutagenesis techniques.

Such variants include those produced by nucleotide substitutions, deletions or additions. The substitutions, deletions or additions may involve one or more nucleotides. These variants may be altered in coding regions, non-coding regions, or both. Alterations in the coding regions may produce conservative or non-conservative amino acid substitutions, deletions or additions. Especially preferred among these are silent substitutions, additions and deletions, which do not alter the properties and activities of the *B. burgdorferi* polypeptides disclosed herein or portions thereof. Also especially preferred in this regard are conservative substitutions.

The present application is further directed to nucleic acid molecules at least 90%, 95%, 96%, 97%, 98% or 99% identical to a nucleic acid sequence shown in Table 1. The above nucleic acid sequences are included irrespective of whether they encode a polypeptide having *B. burgdorferi* activity. This is because even where a particular nucleic acid molecule does not encode a polypeptide having *B. burgdorferi* activity, one of skill in the art would still know how to use the nucleic acid molecule, for instance, as a hybridization probe. Uses of the nucleic acid molecules of the present invention that do not encode a polypeptide having *B. burgdorferi* activity include, *inter alia*, isolating an *B. burgdorferi* gene or allelic variants thereof from a DNA library, and detecting *B. burgdorferi* mRNA expression samples, environmental samples, suspected of containing *B. burgdorferi* by Northern Blot analysis.

Embodiments of the invention include isolated nucleic acid molecules comprising a polynucleotide having a nucleotide sequence at least 90% identical, and more preferably at least 95%, 96%, 97%, 98% or 99% identical to (a) a nucleotide sequence encoding any of the amino acid sequences of the full-length polypeptides shown in Table 1; (b) a nucleotide sequence encoding any of the amino acid sequences of the full-length polypeptides shown in Table 1 but minus the N-terminal methionine residue, if present; (c) a nucleotide sequence encoding any of the

amino acid sequences of the truncated polypeptides shown in Table 1; and (d) a nucleotide sequence complementary to any of the nucleotide sequences in (a), (b), or (c) above.

Preferred, are nucleic acid molecules having sequences at least 90%, 95%, 96%, 97%, 98% or 99% identical to the nucleic acid sequence shown in Table 1, which do, in fact, encode a polypeptide having *B. burgdorferi* protein activity. By "a polypeptide having *B. burgdorferi* activity" is intended polypeptides exhibiting activity similar, but not necessarily identical, to an activity of the *B. burgdorferi* protein of the invention, as measured in a particular biological assay suitable for measuring activity of the specified protein.

Due to the degeneracy of the genetic code, one of ordinary skill in the art will immediately recognize that a large number of the nucleic acid molecules having a sequence at least 90%, 95%, 96%, 97%, 98%, or 99% identical to the nucleic acid sequences shown in Table 1 will encode a polypeptide having *B. burgdorferi* protein activity. In fact, since degenerate variants of these nucleotide sequences all encode the same polypeptide, this will be clear to the skilled artisan even without performing the above described comparison assay. It will be further recognized in the art that, for such nucleic acid molecules that are not degenerate variants, a reasonable number will also encode a polypeptide having *B. burgdorferi* protein activity. This is because the skilled artisan is fully aware of amino acid substitutions that are either less likely or not likely to significantly effect protein function (e.g., replacing one aliphatic amino acid with a second aliphatic amino acid), as further described below.

The biological activity or function of the polypeptides of the present invention are expected to be similar or identical to polypeptides from other bacteria that share a high degree of structural identity/similarity. Tables 2 lists accession numbers and descriptions for the closest matching sequences of polypeptides available through Genbank and Derwent databases. It is therefore expected that the biological activity or function of the polypeptides of the present invention will be similar or identical to those polypeptides from other bacterial genera, species, or strains listed in Table 2.

By a polynucleotide having a nucleotide sequence at least, for example, 95% "identical" to a reference nucleotide sequence of the present invention, it is intended that the nucleotide sequence of the polynucleotide is identical to the reference sequence except that the polynucleotide sequence may include up to five point mutations per each 100 nucleotides of the reference nucleotide sequence encoding the *B. burgdorferi* polypeptide. In other words, to obtain a polynucleotide having a nucleotide sequence at least 95% identical to a reference nucleotide sequence, up to 5% (5 of 100) of the nucleotides in the reference sequence may be deleted, inserted, or substituted with another nucleotide. The query sequence may be an entire sequence shown in Table 1, the ORF (open reading frame), or any fragment specified as described herein.

As a practical matter, whether any particular nucleic acid molecule or polypeptide is at least 90%, 95%, 96%, 97%, 98% or 99% identical to a nucleotide sequence of the present invention can be determined conventionally using known computer programs. A preferred method for determining the best overall match between a query sequence (a sequence of the present invention)

and a subject sequence, also referred to as a global sequence alignment, can be determined using the FASTDB computer program based on the algorithm of Brutlag et al. See Brutlag et al.

(1990) Comp. App. Biosci. 6:237-245. In a sequence alignment the query and subject sequences are both DNA sequences. An RNA sequence can be compared by first converting U's to T's.

5 The result of said global sequence alignment is in percent identity. Preferred parameters used in a FASTDB alignment of DNA sequences to calculate percent identity are: Matrix=Unitary, k-tuple=4, Mismatch Penalty=1, Joining Penalty=30, Randomization Group Length=0, Cutoff Score=1, Gap Penalty=5, Gap Size Penalty 0.05, Window Size=500 or the length of the subject nucleotide sequence, whichever is shorter.

10 If the subject sequence is shorter than the query sequence because of 5' or 3' deletions, not because of internal deletions, a manual correction must be made to the results. This is because the FASTDB program does not account for 5' and 3' truncations of the subject sequence when calculating percent identity. For subject sequences truncated at the 5' or 3' ends, relative to the query sequence, the percent identity is corrected by calculating the number of bases of the query sequence that are 5' and 3' of the subject sequence, which are not matched/aligned, as a percent of the total bases of the query sequence. Whether a nucleotide is matched/aligned is determined by results of the FASTDB sequence alignment. This percentage is then subtracted from the percent identity, calculated by the above FASTDB program using the specified parameters, to arrive at a final percent identity score. This corrected score is what is used for the purposes of the present invention. Only nucleotides outside the 5' and 3' nucleotides of the subject sequence, as displayed by the FASTDB alignment, which are not matched/aligned with the query sequence, are calculated for the purposes of manually adjusting the percent identity score.

25 For example, a 90 nucleotide subject sequence is aligned to a 100 nucleotide query sequence to determine percent identity. The deletions occur at the 5' end of the subject sequence and therefore, the FASTDB alignment does not show a matched/alignment of the first 10 nucleotides at 5' end. The 10 unpaired nucleotides represent 10% of the sequence (number of nucleotides at the 5' and 3' ends not matched/total number of nucleotides in the query sequence) so 10% is subtracted from the percent identity score calculated by the FASTDB program. If the remaining 90 nucleotides were perfectly matched the final percent identity would be 90%. In 30 another example, a 90 nucleotide subject sequence is compared with a 100 nucleotide query sequence. This time the deletions are internal deletions so that there are no nucleotides on the 5' or 3' of the subject sequence which are not matched/aligned with the query. In this case the percent identity calculated by FASTDB is not manually corrected. Once again, only nucleotides 5' and 3' of the subject sequence which are not matched/aligned with the query sequence are 35 manually corrected for. No other manual corrections are to be made for the purposes of the present invention.

Vectors and Host Cells

The present invention also relates to vectors which include the isolated DNA molecules of

above-described host cells are known in the art.

Among vectors preferred for use in bacteria include pQE70, pQE60 and pQE-9, available from Qiagen; pBS vectors, Phagescript vectors, Bluescript vectors, pNH8A, pNH16a, pNH18A, pNH46A available from Stratagene; pET series of vectors available from Novagen; and ptrc99a, pKK223-3, pKK233-3, pDR540, pRIT5 available from Pharmacia. Among preferred eukaryotic vectors are pWLNEO, pSV2CAT, pOG44, pXT1 and pSG available from Stratagene; and pSVK3, pBPV, pMSG and pSVL available from Pharmacia. Other suitable vectors will be readily apparent to the skilled artisan.

Among known bacterial promoters suitable for use in the present invention include the *E. coli lacI* and *lacZ* promoters, the T3 and T7 promoters, the *gpt* promoter, the lambda PR and PL promoters and the *trp* promoter. Suitable eukaryotic promoters include the CMV immediate early promoter, the HSV thymidine kinase promoter, the early and late SV40 promoters, the promoters of retroviral LTRs, such as those of the Rous sarcoma virus (RSV), and metallothionein promoters, such as the mouse metallothionein-I promoter.

Introduction of the construct into the host cell can be effected by calcium phosphate transfection, DEAE-dextran mediated transfection, cationic lipid-mediated transfection, electroporation, transduction, infection or other methods. Such methods are described in many standard laboratory manuals, such as Davis *et al.*, *Basic Methods In Molecular Biology* (1986).

Transcription of DNA encoding the polypeptides of the present invention by higher eukaryotes may be increased by inserting an enhancer sequence into the vector. Enhancers are *cis*-acting elements of DNA, usually about from 10 to 300 bp that act to increase transcriptional activity of a promoter in a given host cell-type. Examples of enhancers include the SV40 enhancer, which is located on the late side of the replication origin at bp 100 to 270, the cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers.

For secretion of the translated polypeptide into the lumen of the endoplasmic reticulum, into the periplasmic space or into the extracellular environment, appropriate secretion signals may be incorporated into the expressed polypeptide. The signals may be endogenous to the polypeptide or they may be heterologous signals.

The polypeptide may be expressed in a modified form, such as a fusion protein, and may include not only secretion signals, but also additional heterologous functional regions. For instance, a region of additional amino acids, particularly charged amino acids, may be added to the N-terminus of the polypeptide to improve stability and persistence in the host cell, during purification, or during subsequent handling and storage. Also, peptide moieties may be added to the polypeptide to facilitate purification. Such regions may be removed prior to final preparation of the polypeptide. The addition of peptide moieties to polypeptides to engender secretion or excretion, to improve stability and to facilitate purification, among others, are familiar and routine techniques in the art. A preferred fusion protein comprises a heterologous region from immunoglobulin that is useful to solubilize proteins. For example, EP-A-O 464 533 (Canadian

the present invention, host cells which are genetically engineered with the recombinant vectors, and the production of *B. burgdorferi* polypeptides or fragments thereof by recombinant techniques.

Recombinant constructs may be introduced into host cells using well known techniques such as infection, transduction, transfection, transvection, electroporation and transformation. The vector may be, for example, a phage, plasmid, viral or retroviral vector. Retroviral vectors may be replication competent or replication defective. In the latter case, viral propagation generally will occur only in complementing host cells.

The polynucleotides may be joined to a vector containing a selectable marker for propagation in a host. Generally, a plasmid vector is introduced in a precipitate, such as a calcium phosphate precipitate, or in a complex with a charged lipid. If the vector is a virus, it may be packaged *in vitro* using an appropriate packaging cell line and then transduced into host cells.

Preferred are vectors comprising *cis*-acting control regions to the polynucleotide of interest. Appropriate *trans*-acting factors may be supplied by the host, supplied by a complementing vector or supplied by the vector itself upon introduction into the host.

In certain preferred embodiments in this regard, the vectors provide for specific expression, which may be inducible and/or cell type-specific. Particularly preferred among such vectors are those inducible by environmental factors that are easy to manipulate, such as temperature and nutrient additives.

Expression vectors useful in the present invention include chromosomal-, episomal- and virus-derived vectors, *e.g.*, vectors derived from bacterial plasmids, bacteriophage, yeast episomes, yeast chromosomal elements, viruses such as baculoviruses, papova viruses, vaccinia viruses, adenoviruses, fowl pox viruses, pseudorabies viruses and retroviruses, and vectors derived from combinations thereof, such as cosmids and phagemids.

The DNA insert should be operatively linked to an appropriate promoter, such as the phage lambda PL promoter, the *E. coli lac*, *trp* and *tac* promoters, the SV40 early and late promoters and promoters of retroviral LTRs, to name a few. Other suitable promoters will be known to the skilled artisan. The expression constructs will further contain sites for transcription initiation, termination and, in the transcribed region, a ribosome binding site for translation. The coding portion of the mature transcripts expressed by the constructs will preferably include a translation initiating site at the beginning and a termination codon (UAA, UGA or UAG) appropriately positioned at the end of the polypeptide to be translated.

As indicated, the expression vectors will preferably include at least one selectable marker. Such markers include dihydrofolate reductase or neomycin resistance for eukaryotic cell culture and tetracycline or ampicillin resistance genes for culturing in *E. coli* and other bacteria. Representative examples of appropriate hosts include, but are not limited to, bacterial cells, such as *E. coli*, *Streptomyces* and *Salmonella typhimurium* cells; fungal cells, such as yeast cells; insect cells such as *Drosophila S2* and *Spodoptera Sf9* cells; animal cells such as CHO, COS and Bowes melanoma cells; and plant cells. Appropriate culture mediums and conditions for the

counterpart 2045869) discloses fusion proteins comprising various portions of constant region of immunoglobulin molecules together with another human protein or part thereof. In many cases, the Fc part in a fusion protein is thoroughly advantageous for use in therapy and diagnosis and thus results, for example, in improved pharmacokinetic properties (EP-A 0232 262). On the other 5 hand, for some uses it would be desirable to be able to delete the Fc part after the fusion protein has been expressed, detected and purified in the advantageous manner described. This is the case when Fc portion proves to be a hindrance to use in therapy and diagnosis, for example when the fusion protein is to be used as antigen for immunizations. In drug discovery, for example, human proteins, such as, hIL5-receptor has been fused with Fc portions for the purpose of 10 high-throughput screening assays to identify antagonists of hIL-5. See Bennett, D. *et al.*, *J. Molec. Recogn.* 8:52-58 (1995) and Johanson, K. *et al.*, *J. Biol. Chem.* 270 (16):9459-9471 (1995).

The *B. burgdorferi* polypeptides can be recovered and purified from recombinant cell cultures by well-known methods including ammonium sulfate or ethanol precipitation, acid extraction, anion or cation exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, affinity chromatography, hydroxylapatite chromatography, lectin chromatography and high performance liquid chromatography ("HPLC") is employed for purification. Polypeptides of the present invention include naturally purified products, products of chemical synthetic procedures, and products produced by recombinant techniques from a prokaryotic or eukaryotic host, including, for example, bacterial, yeast, higher plant, insect and mammalian cells.

Polypeptides and Fragments

The invention further provides isolated polypeptides having the amino acid sequences in 25 Table 1, and peptides or polypeptides comprising portions of the above polypeptides. The terms "peptide" and "oligopeptide" are considered synonymous (as is commonly recognized) and each term can be used interchangeably as the context requires to indicate a chain of at least two amino acids coupled by peptidyl linkages. The word "polypeptide" is used herein for chains containing 30 more than ten amino acid residues. All oligopeptide and polypeptide formulas or sequences herein are written from left to right and in the direction from amino terminus to carboxy terminus.

As discussed in detail below, immunization using *B. burgdorferi* sensu stricto isolate B31 decorin-binding protein elicits the production of antiserum which confers passive immunity against *Borrelia* species and strains which express divergent forms of this protein. Cassatt, D. *et al.*, *Protection of Borrelia burgdorferi Infection by Antibodies to Decorin-binding Protein*, in 35 VACCINES97, Cold Spring Harbor Press (1997), pages 191-195. Thus, some amino acid sequences of the *B. burgdorferi* polypeptides shown in Table 1 can be varied without significantly effecting the antigenicity of the polypeptides. If such differences in sequence are contemplated, it should be remembered that there will be critical areas on the polypeptide which determine antigenicity. In general, it is possible to replace residues which do not form part of an

antigenic epitope without significantly effecting the antigenicity of a polypeptide.

Variant and Mutant Polypeptides

To improve or alter the characteristics of *B. burgdorferi* polypeptides of the present invention, protein engineering may be employed. Recombinant DNA technology known to those skilled in the art can be used to create novel mutant proteins or muteins including single or multiple amino acid substitutions, deletions, additions, or fusion proteins. Such modified polypeptides can show, e.g., enhanced activity or increased stability. In addition, they may be purified in higher yields and show better solubility than the corresponding natural polypeptide, at least under certain purification and storage conditions.

10

***N*-Terminal and *C*-Terminal Deletion Mutants**

It is known in the art that one or more amino acids may be deleted from the N-terminus or C-terminus without substantial loss of biological function. For instance, Ron et al. *J. Biol. Chem.*, 268:2984-2988 (1993), reported modified KGF proteins that had heparin binding activity even if 3, 8, or 27 N-terminal amino acid residues were missing. Accordingly, the present invention provides polypeptides having one or more residues deleted from the amino terminus of the amino acid sequence of the *B. burgdorferi* polypeptides shown in Table 1, and polynucleotides encoding such polypeptides.

Similarly, many examples of biologically functional C-terminal deletion muteins are known. For instance, Interferon gamma shows up to ten times higher activities by deleting 8-10 amino acid residues from the carboxy terminus of the protein *See, e.g., Dobeli, et al. (1988) J. Biotechnology 7:199-216.* Accordingly, the present invention provides polypeptides having one or more residues from the carboxy terminus of the amino acid sequence of the *B. burgdorferi* polypeptides shown in Table 1. The invention also provides polypeptides having one or more amino acids deleted from both the amino and the carboxyl termini as described below.

The present invention is further directed to polynucleotide encoding portions or fragments of the amino acid sequences described herein as well as to portions or fragments of the isolated amino acid sequences described herein. Fragments include portions of the amino acid sequences of Table 1, are at least 5 contiguous amino acid in length, are selected from any two integers, one of which representing a N-terminal position. The initiation codon of the polypeptides of the present invention is position 1. Every combination of a N-terminal and C-terminal position that a fragment at least 5 contiguous amino acid residues in length could occupy, on any given amino acid sequence of Table 1 is included in the invention. At least means a fragment may be 5 contiguous amino acid residues in length or any integer between 5 and the number of residues in a full length amino acid sequence minus 1. Therefore, included in the invention are contiguous fragments specified by any N-terminal and C-terminal positions of amino acid sequence set forth in Table 1 wherein the contiguous fragment is any integer between 5 and the number of residues in a full length sequence minus 1.

Further, the invention includes polypeptides comprising fragments specified by size, in

amino acid residues, rather than by N-terminal and C-terminal positions. The invention includes any fragment size, in contiguous amino acid residues, selected from integers between 5 and the number of residues in a full length sequence minus 1. Preferred sizes of contiguous polypeptide fragments include about 5 amino acid residues, about 10 amino acid residues, about 20 amino acid residues, about 30 amino acid residues, about 40 amino acid residues, about 50 amino acid residues, about 100 amino acid residues, about 200 amino acid residues, about 300 amino acid residues, and about 400 amino acid residues. The preferred sizes are, of course, meant to exemplify, not limit, the present invention as all size fragments representing any integer between 5 and the number of residues in a full length sequence minus 1 are included in the invention. The present invention also provides for the exclusion of any fragments specified by N-terminal and C-terminal positions or by size in amino acid residues as described above. Any number of fragments specified by N-terminal and C-terminal positions or by size in amino acid residues as described above may be excluded.

The above fragments need not be active since they would be useful, for example, in immunoassays, in epitope mapping, epitope tagging, to generate antibodies to a particular portion of the protein, as vaccines, and as molecular weight markers.

Other Mutants

In addition to N- and C-terminal deletion forms of the protein discussed above, it also will be recognized by one of ordinary skill in the art that some amino acid sequences of the *B. burgdorferi* polypeptide can be varied without significant effect of the structure or function of the protein. If such differences in sequence are contemplated, it should be remembered that there will be critical areas on the protein which determine activity.

Thus, the invention further includes variations of the *B. burgdorferi* polypeptides which show substantial *B. burgdorferi* polypeptide activity or which include regions of *B. burgdorferi* protein such as the protein portions discussed below. Such mutants include deletions, insertions, inversions, repeats, and type substitutions selected according to general rules known in the art so as to have little effect on activity. For example, guidance concerning how to make phenotypically silent amino acid substitutions is provided. There are two main approaches for studying the tolerance of an amino acid sequence to change. *See*, Bowie, J. U. *et al.* (1990), *Science* 247:1306-1310. The first method relies on the process of evolution, in which mutations are either accepted or rejected by natural selection. The second approach uses genetic engineering to introduce amino acid changes at specific positions of a cloned gene and selections or screens to identify sequences that maintain functionality.

These studies have revealed that proteins are surprisingly tolerant of amino acid substitutions. The studies indicate which amino acid changes are likely to be permissive at a certain position of the protein. For example, most buried amino acid residues require nonpolar side chains, whereas few features of surface side chains are generally conserved. Other such phenotypically silent substitutions are described by Bowie *et al.* (*supra*) and the references cited

therein. Typically seen as conservative substitutions are the replacements, one for another, among the aliphatic amino acids Ala, Val, Leu and Ile; interchange of the hydroxyl residues Ser and Thr, exchange of the acidic residues Asp and Glu, substitution between the amide residues Asn and Gln, exchange of the basic residues Lys and Arg and replacements among the aromatic residues Phe, Tyr.

Thus, the fragment, derivative, analog, or homolog of the polypeptide of Table 1, or that encoded by the plamids listed in Table 1, may be: (i) one in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue) and such substituted amino acid residue may or may not be one encoded by the genetic code; or (ii) one in which one or more of the amino acid residues includes a substituent group; or (iii) one in which the *B. burgdorferi* polypeptide is fused with another compound, such as a compound to increase the half-life of the polypeptide (for example, polyethylene glycol); or (iv) one in which the additional amino acids are fused to the above form of the polypeptide, such as an IgG Fc fusion region peptide or leader or secretory sequence or a sequence which is employed for purification of the above form of the polypeptide or a proprotein sequence. Such fragments, derivatives and analogs are deemed to be within the scope of those skilled in the art from the teachings herein.

Thus, the *B. burgdorferi* polypeptides of the present invention may include one or more amino acid substitutions, deletions, or additions, either from natural mutations or human manipulation. As indicated, changes are preferably of a minor nature, such as conservative amino acid substitutions that do not significantly affect the folding or activity of the protein (see Table 3).

Amino acids in the *B. burgdorferi* proteins of the present invention that are essential for function can be identified by methods known in the art, such as site-directed mutagenesis or alanine-scanning mutagenesis. *See, e.g.*, Cunningham et al. (1989) *Science* 244:1081-1085. The latter procedure introduces single alanine mutations at every residue in the molecule. The resulting mutant molecules are then tested for biological activity using assays appropriate for measuring the function of the particular protein.

Of special interest are substitutions of charged amino acids with other charged or neutral amino acids which may produce proteins with highly desirable improved characteristics, such as less aggregation. Aggregation may not only reduce activity but also be problematic when preparing pharmaceutical formulations, because aggregates can be immunogenic. *See, e.g.*, Pinckard et al., (1967) *Clin. Exp. Immunol.* 2:331-340; Robbins, et al., (1987) *Diabetes* 36:838-845; Cleland, et al., (1993) *Crit. Rev. Therapeutic Drug Carrier Systems* 10:307-377.

The polypeptides of the present invention are preferably provided in an isolated form, and preferably are substantially purified. A recombinantly produced version of the *B. burgdorferi* polypeptide can be substantially purified by the one-step method described by Smith et al. (1988) *Gene* 67:31-40. Polypeptides of the invention also can be purified from natural or recombinant sources using antibodies directed against the polypeptides of the invention in methods which are well known in the art of protein purification.

The invention further provides for isolated *B. burgdorferi* polypeptides comprising an amino acid sequence selected from the group consisting of: (a) the amino acid sequence of a full-length *B. burgdorferi* polypeptide having the complete amino acid sequence shown in Table 1; (b) the amino acid sequence of a full-length *B. burgdorferi* polypeptide having the complete amino acid sequence shown in Table 1 excepting the N-terminal methionine; (c) the complete amino acid sequence encoded by the plamids listed in Table 1; and (d) the complete amino acid sequence excepting the N-terminal methionine encoded by the plamids listed in Table 1. The polypeptides of the present invention also include polypeptides having an amino acid sequence at least 80% identical, more preferably at least 90% identical, and still more preferably 95%, 96%, 97%, 98% or 99% identical to those described in (a), (b), (c), and (d) above.

Further polypeptides of the present invention include polypeptides which have at least 90% similarity, more preferably at least 95% similarity, and still more preferably at least 96%, 97%, 98% or 99% similarity to those described above.

A further embodiment of the invention relates to a polypeptide which comprises the amino acid sequence of a *B. burgdorferi* polypeptide having an amino acid sequence which contains at least one conservative amino acid substitution, but not more than 50 conservative amino acid substitutions, not more than 40 conservative amino acid substitutions, not more than 30 conservative amino acid substitutions, and not more than 20 conservative amino acid substitutions. Also provided are polypeptides which comprise the amino acid sequence of a *B. burgdorferi* polypeptide, having at least one, but not more than 10, 9, 8, 7, 6, 5, 4, 3, 2 or 1 conservative amino acid substitutions.

By a polypeptide having an amino acid sequence at least, for example, 95% "identical" to a query amino acid sequence of the present invention, it is intended that the amino acid sequence of the subject polypeptide is identical to the query sequence except that the subject polypeptide sequence may include up to five amino acid alterations per each 100 amino acids of the query amino acid sequence. In other words, to obtain a polypeptide having an amino acid sequence at least 95% identical to a query amino acid sequence, up to 5% of the amino acid residues in the subject sequence may be inserted, deleted, (indels) or substituted with another amino acid. These alterations of the reference sequence may occur at the amino or carboxy terminal positions of the reference amino acid sequence or anywhere between those terminal positions, interspersed either individually among residues in the reference sequence or in one or more contiguous groups within the reference sequence.

As a practical matter, whether any particular polypeptide is at least 90%, 95%, 96%, 97%, 98% or 99% identical to, for instance, the amino acid sequences shown in Table 1 or to the amino acid sequence encoded by the plamids listed in Table 1 can be determined conventionally using known computer programs. A preferred method for determining the best overall match between a query sequence (a sequence of the present invention) and a subject sequence, also referred to as a global sequence alignment, can be determined using the FASTDB computer program based on the algorithm of Brutlag et al., (1990) Comp. App. Biosci. 6:237-245. In a sequence alignment the

query and subject sequences are both amino acid sequences. The result of said global sequence alignment is in percent identity. Preferred parameters used in a FASTDB amino acid alignment are: Matrix=PAM 0, k-tuple=2, Mismatch Penalty=1, Joining Penalty=20, Randomization Group Length=0, Cutoff Score=1, Window Size=sequence length, Gap Penalty=5, Gap Size 5 Penalty=0.05, Window Size=500 or the length of the subject amino acid sequence, whichever is shorter.

If the subject sequence is shorter than the query sequence due to N- or C-terminal deletions, not because of internal deletions, the results, in percent identity, must be manually corrected. This is because the FASTDB program does not account for N- and C-terminal truncations of the subject sequence when calculating global percent identity. For subject sequences truncated at the N- and C-termini, relative to the query sequence, the percent identity is corrected by calculating the number of residues of the query sequence that are N- and C-terminal of the subject sequence, which are not matched/aligned with a corresponding subject residue, as a percent of the total bases of the query sequence. Whether a residue is matched/aligned is determined by results of the FASTDB sequence alignment. This percentage is then subtracted from the percent identity, calculated by the above FASTDB program using the specified parameters, to arrive at a final percent identity score. This final percent identity score is what is used for the purposes of the present invention. Only residues to the N- and C-termini of the subject sequence, which are not matched/aligned with the query sequence, are considered for the purposes of manually adjusting the percent identity score. That is, only query amino acid residues outside the farthest N- and C-terminal residues of the subject sequence.

For example, a 90 amino acid residue subject sequence is aligned with a 100 residue query sequence to determine percent identity. The deletion occurs at the N-terminus of the subject sequence and therefore, the FASTDB alignment does not match/align with the first 10 residues at the N-terminus. The 10 unpaired residues represent 10% of the sequence (number of residues at the N- and C- termini not matched/total number of residues in the query sequence) so 10% is subtracted from the percent identity score calculated by the FASTDB program. If the remaining 90 residues were perfectly matched the final percent identity would be 90%. In another example, a 90 residue subject sequence is compared with a 100 residue query sequence. This time the 25 deletions are internal so there are no residues at the N- or C-termini of the subject sequence which are not matched/aligned with the query. In this case the percent identity calculated by FASTDB is not manually corrected. Once again, only residue positions outside the N- and C-terminal ends of the subject sequence, as displayed in the FASTDB alignment, which are not matched/aligned 30 with the query sequence are manually corrected. No other manual corrections are to be made for the purposes of the present invention.

The above polypeptide sequences are included irrespective of whether they have their normal biological activity. This is because even where a particular polypeptide molecule does not have biological activity, one of skill in the art would still know how to use the polypeptide, for instance, as a vaccine or to generate antibodies. Other uses of the polypeptides of the present

invention that do not have *B. burgdorferi* activity include, *inter alia*, as epitope tags, in epitope mapping, and as molecular weight markers on SDS-PAGE gels or on molecular sieve gel filtration columns using methods known to those of skill in the art.

As described below, the polypeptides of the present invention can also be used to raise 5 polyclonal and monoclonal antibodies, which are useful in assays for detecting *B. burgdorferi* protein expression or as agonists and antagonists capable of enhancing or inhibiting *B. burgdorferi* protein function. Further, such polypeptides can be used in the yeast two-hybrid system to "capture" *B. burgdorferi* protein binding proteins which are also candidate agonists and antagonists according to the present invention. *See, e.g.*, Fields et al. (1989) *Nature* 10 340:245-246.

Epitope-Bearing Portions

In another aspect, the invention provides peptides and polypeptides comprising epitope-bearing portions of the *B. burgdorferi* polypeptides of the present invention. These epitopes are immunogenic or antigenic epitopes of the polypeptides of the present invention. An "immunogenic epitope" is defined as a part of a protein that elicits an antibody response when the whole protein or polypeptide is the immunogen. These immunogenic epitopes are believed to be confined to a few loci on the molecule. On the other hand, a region of a protein molecule to which an antibody can bind is defined as an "antigenic determinant" or "antigenic epitope." The number of immunogenic epitopes of a protein generally is less than the number of antigenic epitopes. *See, e.g.*, Geysen, et al. (1983) *Proc. Natl. Acad. Sci. USA* 81:3998-4002. Predicted antigenic epitopes are shown in Table 4, below. It is pointed out that Table 4 only lists 25 amino acid residues comprising epitopes predicted to have the highest degree of antigenicity. The polypeptides not listed in Table 4 and portions of polypeptides not listed in Table 4 are not considered non-antigenic. This is because they may still be antigenic *in vivo* but merely not 30 recognized as such by the particular algorithm used. Thus, Table 4 lists the amino acid residues comprising preferred antigenic epitopes but not a complete list. Amino acid residues comprising other antigenic epitopes may be determined by algorithms similar to the Jameson-Wolf analysis or by *in vivo* testing for an antigenic response using the methods described herein or those known in the art.

As to the selection of peptides or polypeptides bearing an antigenic epitope (*i.e.*, that contain a region of a protein molecule to which an antibody can bind), it is well known in that art that relatively short synthetic peptides that mimic part of a protein sequence are routinely capable of eliciting an antiserum that reacts with the partially mimicked protein. *See, e.g.*, Sutcliffe, et 35 al., (1983) *Science* 219:660-666. Peptides capable of eliciting protein-reactive sera are frequently represented in the primary sequence of a protein, can be characterized by a set of simple chemical rules, and are confined neither to immunodominant regions of intact proteins (*i.e.*, immunogenic epitopes) nor to the amino or carboxyl terminals. Peptides that are extremely hydrophobic and those of six or fewer residues generally are ineffective at inducing antibodies that bind to the

mimicked protein; longer, peptides, especially those containing proline residues, usually are effective. *See, Sutcliffe, et al., supra, p. 661.* For instance, 18 of 20 peptides designed according to these guidelines, containing 8-39 residues covering 75% of the sequence of the influenza virus hemagglutinin HA1 polypeptide chain, induced antibodies that reacted with the HA1 protein or intact virus; and 12/12 peptides from the MuLV polymerase and 18/18 from the rabies glycoprotein induced antibodies that precipitated the respective proteins.

Antigenic epitope-bearing peptides and polypeptides of the invention are therefore useful to raise antibodies, including monoclonal antibodies, that bind specifically to a polypeptide of the invention. Thus, a high proportion of hybridomas obtained by fusion of spleen cells from donors immunized with an antigen epitope-bearing peptide generally secrete antibody reactive with the native protein. *See Sutcliffe, et al., supra, p. 663.* The antibodies raised by antigenic epitope-bearing peptides or polypeptides are useful to detect the mimicked protein, and antibodies to different peptides may be used for tracking the fate of various regions of a protein precursor which undergoes post-translational processing. The peptides and anti-peptide antibodies may be used in a variety of qualitative or quantitative assays for the mimicked protein, for instance in competition assays since it has been shown that even short peptides (e.g., about 9 amino acids) can bind and displace the larger peptides in immunoprecipitation assays. *See, e.g., Wilson, et al., (1984) Cell 37:767-778.* The anti-peptide antibodies of the invention also are useful for purification of the mimicked protein, for instance, by adsorption chromatography using methods known in the art.

Antigenic epitope-bearing peptides and polypeptides of the invention designed according to the above guidelines preferably contain a sequence of at least seven, more preferably at least nine and most preferably between about 10 to about 50 amino acids (i.e. any integer between 7 and 50) contained within the amino acid sequence of a polypeptide of the invention. However, peptides or polypeptides comprising a larger portion of an amino acid sequence of a polypeptide of the invention, containing about 50 to about 100 amino acids, or any length up to and including the entire amino acid sequence of a polypeptide of the invention, also are considered epitope-bearing peptides or polypeptides of the invention and also are useful for inducing antibodies that react with the mimicked protein. Preferably, the amino acid sequence of the epitope-bearing peptide is selected to provide substantial solubility in aqueous solvents (i.e., the sequence includes relatively hydrophilic residues and highly hydrophobic sequences are preferably avoided); and sequences containing proline residues are particularly preferred.

Non-limiting examples of antigenic polypeptides or peptides that can be used to generate an *Borrelia*-specific immune response or antibodies include portions of the amino acid sequences identified in Table 1. More specifically, Table 4 discloses a list of non-limiting residues that are involved in the antigenicity of the epitope-bearing fragments of the present invention. Therefore, the present inventions provides for isolated and purified antigenic epitope-bearing fragments of the polypeptides of the present invention comprising a peptide sequences of Table 4. The antigenic epitope-bearing fragments comprising a peptide sequence of Table 4 preferably contain a

sequence of at least seven, more preferably at least nine and most preferably between about 10 to about 50 amino acids (i.e. any integer between 7 and 50) of a polypeptide of the present invention. That is, included in the present invention are antigenic polypeptides between the integers of 7 and 50 amino acid in length comprising one or more of the sequences of Table 4.

5 Therefore, in most cases, the polypeptides of Table 4 make up only a portion of the antigenic polypeptide. All combinations of sequences between the integers of 7 and 50 amino acid in length comprising one or more of the sequences of Table 4 are included. The antigenic epitope-bearing fragements may be specified by either the number of contiguous amino acid residues or by specific N-terminal and C-terminal positions as described above for the polypeptide fragements of
10 the present invention, wherein the initiation codon is residue 1. Any number of the described antigenic epitope-bearing fragements of the present invention may also be excluded from the present invention in the same manner.

15 The epitope-bearing peptides and polypeptides of the invention may be produced by any conventional means for making peptides or polypeptides including recombinant means using nucleic acid molecules of the invention. For instance, an epitope-bearing amino acid sequence of the present invention may be fused to a larger polypeptide which acts as a carrier during recombinant production and purification, as well as during immunization to produce anti-peptide antibodies. Epitope-bearing peptides also may be synthesized using known methods of chemical synthesis. For instance, Houghten has described a simple method for synthesis of large numbers of peptides, such as 10-20 mg of 248 different 13 residue peptides representing single amino acid variants of a segment of the HA1 polypeptide which were prepared and characterized (by
20 ELISA-type binding studies) in less than four weeks (Houghten, R. A. Proc. Natl. Acad. Sci. USA 82:5131-5135 (1985)). This "Simultaneous Multiple Peptide Synthesis (SMPS)" process is further described in U.S. Patent No. 4,631,211 to Houghten and coworkers (1986). In this
25 procedure the individual resins for the solid-phase synthesis of various peptides are contained in separate solvent-permeable packets, enabling the optimal use of the many identical repetitive steps involved in solid-phase methods. A completely manual procedure allows 500-1000 or more syntheses to be conducted simultaneously (Houghten et al. (1985) Proc. Natl. Acad. Sci. 82:5131-5135 at 5134).

30 Epitope-bearing peptides and polypeptides of the invention are used to induce antibodies according to methods well known in the art. *See, e.g.*, Sutcliffe, et al., *supra*; Wilson, et al., *supra*; and Bittle, et al. (1985) J. Gen. Virol. 66:2347-2354. Generally, animals may be immunized with free peptide; however, anti-peptide antibody titer may be boosted by coupling of the peptide to a macromolecular carrier, such as keyhole limpet hemacyanin (KLH) or tetanus
35 toxoid. For instance, peptides containing cysteine may be coupled to carrier using a linker such as m-maleimidobenzoyl-N-hydroxysuccinimide ester (MBS), while other peptides may be coupled to carrier using a more general linking agent such as glutaraldehyde. Animals such as rabbits, rats and mice are immunized with either free or carrier-coupled peptides, for instance, by intraperitoneal and/or intradermal injection of emulsions containing about 100 µg peptide or

carrier protein and Freund's adjuvant. Several booster injections may be needed, for instance, at intervals of about two weeks, to provide a useful titer of anti-peptide antibody which can be detected, for example, by ELISA assay using free peptide adsorbed to a solid surface. The titer of anti-peptide antibodies in serum from an immunized animal may be increased by selection of anti-peptide antibodies, for instance, by adsorption to the peptide on a solid support and elution of the selected antibodies according to methods well known in the art.

5 Immunogenic epitope-bearing peptides of the invention, *i.e.*, those parts of a protein that elicit an antibody response when the whole protein is the immunogen, are identified according to methods known in the art. For instance, Geysen, *et al.*, *supra*, discloses a procedure for rapid 10 concurrent synthesis on solid supports of hundreds of peptides of sufficient purity to react in an ELISA. Interaction of synthesized peptides with antibodies is then easily detected without removing them from the support. In this manner a peptide bearing an immunogenic epitope of a desired protein may be identified routinely by one of ordinary skill in the art. For instance, the immunologically important epitope in the coat protein of foot-and-mouth disease virus was located by Geysen *et al.* *supra* with a resolution of seven amino acids by synthesis of an overlapping set 15 of all 208 possible hexapeptides covering the entire 213 amino acid sequence of the protein. Then, a complete replacement set of peptides in which all 20 amino acids were substituted in turn at every position within the epitope were synthesized, and the particular amino acids conferring 20 specificity for the reaction with antibody were determined. Thus, peptide analogs of the epitope-bearing peptides of the invention can be made routinely by this method. U.S. Patent No. 4,708,781 to Geysen (1987) further describes this method of identifying a peptide bearing an immunogenic epitope of a desired protein.

25 Further still, U.S. Patent No. 5,194,392, to Geysen (1990), describes a general method of detecting or determining the sequence of monomers (amino acids or other compounds) which is a topological equivalent of the epitope (*i.e.*, a "mimotope") which is complementary to a particular paratope (antigen binding site) of an antibody of interest. More generally, U.S. Patent 30 No. 4,433,092, also to Geysen (1989), describes a method of detecting or determining a sequence of monomers which is a topographical equivalent of a ligand which is complementary to the ligand binding site of a particular receptor of interest. Similarly, U.S. Patent No. 5,480,971 to Houghten, R. A. *et al.* (1996) discloses linear C₁-C₇-alkyl peralkylated oligopeptides and sets 35 and libraries of such peptides, as well as methods for using such oligopeptide sets and libraries for determining the sequence of a peralkylated oligopeptide that preferentially binds to an acceptor molecule of interest. Thus, non-peptide analogs of the epitope-bearing peptides of the invention also can be made routinely by these methods. The entire disclosure of each document cited in this section on "Polypeptides and Fragments" is hereby incorporated herein by reference.

As one of skill in the art will appreciate, the polypeptides of the present invention and the epitope-bearing fragments thereof described above can be combined with parts of the constant domain of immunoglobulins (IgG), resulting in chimeric polypeptides. These fusion proteins facilitate purification and show an increased half-life *in vivo*. This has been shown, *e.g.*, for

chimeric proteins consisting of the first two domains of the human CD4-polypeptide and various domains of the constant regions of the heavy or light chains of mammalian immunoglobulins. (EPA 0,394,827; Traunecker et al. (1988) *Nature* 331:84-86. Fusion proteins that have a disulfide-linked dimeric structure due to the IgG part can also be more efficient in binding and neutralizing other molecules than a monomeric *B. burgdorferi* polypeptide or fragment thereof alone. See Fountoulakis et al. (1995) *J. Biochem.* 270:3958-3964. Nucleic acids encoding the above epitopes of *B. burgdorferi* polypeptides can also be recombined with a gene of interest as an epitope tag to aid in detection and purification of the expressed polypeptide.

10 *Antibodies*

15 *B. burgdorferi* protein-specific antibodies for use in the present invention can be raised against the intact *B. burgdorferi* protein or an antigenic polypeptide fragment thereof, which may be presented together with a carrier protein, such as an albumin, to an animal system (such as rabbit or mouse) or, if it is long enough (at least about 25 amino acids), without a carrier.

20 As used herein, the term "antibody" (Ab) or "monoclonal antibody" (Mab) is meant to include intact molecules, single chain whole antibodies, and antibody fragments. Antibody fragments of the present invention include Fab and F(ab')2 and other fragments including single-chain Fvs (scFv) and disulfide-linked Fvs (sdFv). Also included in the present invention are chimeric and humanized monoclonal antibodies and polyclonal antibodies specific for the 25 polypeptides of the present invention. The antibodies of the present invention may be prepared by any of a variety of methods. For example, cells expressing a polypeptide of the present invention or an antigenic fragment thereof can be administered to an animal in order to induce the production of sera containing polyclonal antibodies. For example, a preparation of *B. burgdorferi* polypeptide or fragment thereof is prepared and purified to render it substantially free of natural 30 contaminants. Such a preparation is then introduced into an animal in order to produce polyclonal antisera of greater specific activity.

35 In a preferred method, the antibodies of the present invention are monoclonal antibodies or binding fragments thereof. Such monoclonal antibodies can be prepared using hybridoma technology. See, e.g., Harlow et al., *ANTIBODIES: A LABORATORY MANUAL*, (Cold Spring Harbor Laboratory Press, 2nd ed. 1988); Hammerling, et al., in: *MONOCLONAL ANTIBODIES AND T-CELL HYBRIDOMAS* 563-681 (Elsevier, N.Y., 1981). Fab and F(ab')2 fragments may be produced by proteolytic cleavage, using enzymes such as papain (to produce Fab fragments) or pepsin (to produce F(ab')2 fragments). Alternatively, *B. burgdorferi* polypeptide-binding fragments, chimeric, and humanized antibodies can be produced through the application of recombinant DNA technology or through synthetic chemistry using methods known in the art.

40 Alternatively, additional antibodies capable of binding to the polypeptide antigen of the present invention may be produced in a two-step procedure through the use of anti-idiotypic antibodies. Such a method makes use of the fact that antibodies are themselves antigens, and that,

therefore, it is possible to obtain an antibody which binds to a second antibody. In accordance with this method, *B. burgdorferi* polypeptide-specific antibodies are used to immunize an animal, preferably a mouse. The splenocytes of such an animal are then used to produce hybridoma cells, and the hybridoma cells are screened to identify clones which produce an antibody whose ability to bind to the *B. burgdorferi* polypeptide-specific antibody can be blocked by the *B. burgdorferi* polypeptide antigen. Such antibodies comprise anti-idiotypic antibodies to the *B. burgdorferi* polypeptide-specific antibody and can be used to immunize an animal to induce formation of further *B. burgdorferi* polypeptide-specific antibodies.

Antibodies and fragement thereof of the present invention may be described by the portion of a polypeptide of the present invention recognized or specifically bound by the antibody. Antibody binding fragement of a polypeptide of the present invention may be described or specified in the same manner as for polypeptide framents discussed above., i.e, by N-terminal and C-terminal positions or by size in contiguous amino acid residues. Any number of antibody binding fragments, of a polypeptide of the present invention, specified by N-terminal and C-terminal positions or by size in amino acid residues, as described above, may also be excluded from the present invention. Therefore, the present invention includes antibodies the specifically bind a particullarly described fragement of a polypeptide of the present invention and allows for the exclusion of the same.

Antibodies and framents thereof of the present invention may also be described or specified in terms of their cross-reactivity. Antibodies and framents that do not bind polypeptides of any other species of *Borrelia* other than *B. burgdorferi* are included in the present invention. Likewise, antibodies and framents that bind only species of *Borrelia*, i.e. antibodies and framents that do not bind bacteria from any genus other than *Borrelia*, are included in the present invention.

25 *Diagnostic Assays*

The present invention further relates to methods for assaying *staphylococcal* infection in an animal by detecting the expression of genes encoding *staphylococcal* polypeptides of the present invention. The methods comprise analyzing tissue or body fluid from the animal for *Borrelia*-specific antibodies, nucleic acids, or proteins. Analysis of nucleic acid specific to *Borrelia* is assayed by PCR or hybridization techniques using nucleic acid sequences of the present invention as either hybridization probes or primers. See, e.g., Sambrook et al. Molecular cloning: A Laboratory Manual (Cold Spring Harbor Laboratory Press, 2nd ed., 1989, page 54 reference); Eremeeva et al. (1994) J. Clin. Microbiol. 32:803-810 (describing differentiation among spotted fever group *Rickettsiae* species by analysis of restriction fragment length polymorphism of PCR-amplified DNA) and Chen et al. 1994 J. Clin. Microbiol. 32:589-595 (detecting *B. burgdorferi* nucleic acids via PCR).

Where diagnosis of a disease state related to infection with *Borrelia* has already been made, the present invention is useful for monitoring progression or regression of the disease state

whereby patients exhibiting enhanced *Borrelia* gene expression will experience a worse clinical outcome relative to patients expressing these gene(s) at a lower level.

By "biological sample" is intended any biological sample obtained from an animal, cell line, tissue culture, or other source which contains *Borrelia* polypeptide, mRNA, or DNA.

5 Biological samples include body fluids (such as saliva, blood, plasma, urine, mucus, synovial fluid, etc.) tissues (such as muscle, skin, and cartilage) and any other biological source suspected of containing *Borrelia* polypeptides or nucleic acids. Methods for obtaining biological samples such as tissue are well known in the art.

10 The present invention is useful for detecting diseases related to *Borrelia* infections in animals. Preferred animals include monkeys, apes, cats, dogs, birds, cows, pigs, mice, horses, rabbits and humans. Particularly preferred are humans.

15 Total RNA can be isolated from a biological sample using any suitable technique such as the single-step guanidinium-thiocyanate-phenol-chloroform method described in Chomczynski et al. (1987) *Anal. Biochem.* 162:156-159. mRNA encoding *Borrelia* polypeptides having sufficient homology to the nucleic acid sequences identified in Table 1 to allow for hybridization between complementary sequences are then assayed using any appropriate method. These include Northern blot analysis, S1 nuclease mapping; the polymerase chain reaction (PCR), reverse transcription in combination with the polymerase chain reaction (RT-PCR), and reverse transcription in combination with the ligase chain reaction (RT-LCR).

20 Northern blot analysis can be performed as described in Harada et al. (1990) *Cell* 63:303-312. Briefly, total RNA is prepared from a biological sample as described above. For the Northern blot, the RNA is denatured in an appropriate buffer (such as glyoxal/dimethyl sulfoxide/sodium phosphate buffer), subjected to agarose gel electrophoresis, and transferred onto a nitrocellulose filter. After the RNAs have been linked to the filter by a UV linker, the filter is prehybridized in a solution containing formamide, SSC, Denhardt's solution, denatured salmon sperm, SDS, and sodium phosphate buffer. A *B. burgdorferi* polynucleotide sequence shown in Table 1 labeled according to any appropriate method (such as the ³²P-multiprime DNA labeling system (Amersham)) is used as probe. After hybridization overnight, the filter is washed and exposed to x-ray film. DNA for use as probe according to the present invention is described in the sections above and will preferably at least 15 nucleotides in length.

25 S1 mapping can be performed as described in Fujita et al. (1987) *Cell* 49:357-367. To prepare probe DNA for use in S1 mapping, the sense strand of an above-described *B. burgdorferi* DNA sequence of the present invention is used as a template to synthesize labeled antisense DNA. The antisense DNA can then be digested using an appropriate restriction endonuclease to generate further DNA probes of a desired length. Such antisense probes are useful for visualizing protected bands corresponding to the target mRNA (i.e., mRNA encoding *Borrelia* polypeptides).

30 Levels of mRNA encoding *Borrelia* polypeptides are assayed, for e.g., using the RT-PCR method described in Makino et al. (1990) *Technique* 2:295-301. By this method, the radioactivities of the "amplicons" in the polyacrylamide gel bands are linearly related to the initial

concentration of the target mRNA. Briefly, this method involves adding total RNA isolated from a biological sample in a reaction mixture containing a RT primer and appropriate buffer. After incubating for primer annealing, the mixture can be supplemented with a RT buffer, dNTPs, DTT, RNase inhibitor and reverse transcriptase. After incubation to achieve reverse transcription of the RNA, the RT products are then subject to PCR using labeled primers. Alternatively, rather than labeling the primers, a labeled dNTP can be included in the PCR reaction mixture. PCR amplification can be performed in a DNA thermal cycler according to conventional techniques. After a suitable number of rounds to achieve amplification, the PCR reaction mixture is electrophoresed on a polyacrylamide gel. After drying the gel, the radioactivity of the appropriate bands (corresponding to the mRNA encoding the *Borrelia* polypeptides of the present invention) are quantified using an imaging analyzer. RT and PCR reaction ingredients and conditions, reagent and gel concentrations, and labeling methods are well known in the art. Variations on the RT-PCR method will be apparent to the skilled artisan. Other PCR methods that can detect the nucleic acid of the present invention can be found in PCR PRIMER: A LABORATORY MANUAL (C.W. Dieffenbach et al. eds., Cold Spring Harbor Lab Press, 1995).

The polynucleotides of the present invention, including both DNA and RNA, may be used to detect polynucleotides of the present invention or *Borrelia* species including *B. burgdorferi* using bio chip technology. The present invention includes both high density chip arrays (>1000 oligonucleotides per cm²) and low density chip arrays (<1000 oligonucleotides per cm²). Bio chips comprising arrays of polynucleotides of the present invention may be used to detect *Borrelia* species, including *B. burgdorferi*, in biological and environmental samples and to diagnose an animal, including humans, with an *B. burgdorferi* or other *Borrelia* infection. The bio chips of the present invention may comprise polynucleotide sequences of other pathogens including bacteria, viral, parasitic, and fungal polynucleotide sequences, in addition to the polynucleotide sequences of the present invention, for use in rapid differential pathogenic detection and diagnosis. The bio chips can also be used to monitor an *B. burgdorferi* or other *Borrelia* infections and to monitor the genetic changes (deletions, insertions, mismatches, etc.) in response to drug therapy in the clinic and drug development in the laboratory. The bio chip technology comprising arrays of polynucleotides of the present invention may also be used to simultaneously monitor the expression of a multiplicity of genes, including those of the present invention. The polynucleotides used to comprise a selected array may be specified in the same manner as for the fragments, i.e., by their 5' and 3' positions or length in contiguous base pairs and include from. Methods and particular uses of the polynucleotides of the present invention to detect *Borrelia* species, including *B. burgdorferi*, using bio chip technology include those known in the art and those of: U.S. Patent Nos. 5510270, 5545531, 5445934, 5677195, 5532128, 5556752, 5527681, 5451683, 5424186, 5607646, 5658732 and World Patent Nos. WO/9710365, WO/9511995, WO/9743447, WO/9535505, each incorporated herein in their entireties.

Biosensors using the polynucleotides of the present invention may also be used to detect, diagnose, and monitor *B. burgdorferi* or other *Borrelia* species and infections thereof.

Biosensors using the polynucleotides of the present invention may also be used to detect particular polynucleotides of the present invention. Biosensors using the polynucleotides of the present invention may also be used to monitor the genetic changes (deletions, insertions, mismatches, etc.) in response to drug therapy in the clinic and drug development in the laboratory. Methods and particular uses of the polynucleotides of the present invention to detect *Borrelia* species, including *B. burgdorferi*, using biosensors include those known in the art and those of: U.S. Patent Nos 5721102, 5658732, 5631170, and World Patent Nos. WO97/35011, WO/9720203, each incorporated herein in their entireties.

Thus, the present invention includes both bio chips and biosensors comprising polynucleotides of the present invention and methods of their use.

Assaying *Borrelia* polypeptide levels in a biological sample can occur using any art-known method, such as antibody-based techniques. For example, *Borrelia* polypeptide expression in tissues can be studied with classical immunohistological methods. In these, the specific recognition is provided by the primary antibody (polyclonal or monoclonal) but the secondary detection system can utilize fluorescent, enzyme, or other conjugated secondary antibodies. As a result, an immunohistological staining of tissue section for pathological examination is obtained. Tissues can also be extracted, e.g., with urea and neutral detergent, for the liberation of *Borrelia* polypeptides for Western-blot or dot/slot assay. See, e.g., Jalkanen, M. et al. (1985) J. Cell. Biol. 101:976-985; Jalkanen, M. et al. (1987) J. Cell. Biol. 105:3087-3096. In this technique, which is based on the use of cationic solid phases, quantitation of a *Borrelia* polypeptide can be accomplished using an isolated *Borrelia* polypeptide as a standard. This technique can also be applied to body fluids.

Other antibody-based methods useful for detecting *Borrelia* polypeptide gene expression include immunoassays, such as the ELISA and the radioimmunoassay (RIA). For example, a *Borrelia* polypeptide-specific monoclonal antibodies can be used both as an immunoabsorbent and as an enzyme-labeled probe to detect and quantify a *Borrelia* polypeptide. The amount of a *Borrelia* polypeptide present in the sample can be calculated by reference to the amount present in a standard preparation using a linear regression computer algorithm. Such an ELISA is described in Iacobelli et al. (1988) Breast Cancer Research and Treatment 11:19-30. In another ELISA assay, two distinct specific monoclonal antibodies can be used to detect *Borrelia* polypeptides in a body fluid. In this assay, one of the antibodies is used as the immunoabsorbent and the other as the enzyme-labeled probe.

The above techniques may be conducted essentially as a "one-step" or "two-step" assay. The "one-step" assay involves contacting the *Borrelia* polypeptide with immobilized antibody and, without washing, contacting the mixture with the labeled antibody. The "two-step" assay involves washing before contacting the mixture with the labeled antibody. Other conventional methods may also be employed as suitable. It is usually desirable to immobilize one component of the assay system on a support, thereby allowing other components of the system to be brought into contact with the component and readily removed from the sample. Variations of the above

and other immunological methods included in the present invention can also be found in Harlow et al., *ANTIBODIES: A LABORATORY MANUAL*, (Cold Spring Harbor Laboratory Press, 2nd ed. 1988).

Suitable enzyme labels include, for example, those from the oxidase group, which catalyze the production of hydrogen peroxide by reacting with substrate. Glucose oxidase is particularly preferred as it has good stability and its substrate (glucose) is readily available.

Activity of an oxidase label may be assayed by measuring the concentration of hydrogen peroxide formed by the enzyme-labeled antibody/substrate reaction. Besides enzymes, other suitable labels include radioisotopes, such as iodine (^{125}I , ^{121}I), carbon (^{14}C), sulphur (^{35}S), tritium (^3H), indium (^{112}In), and technetium ($^{99\text{m}}\text{Tc}$), and fluorescent labels, such as fluorescein and rhodamine, and biotin.

Further suitable labels for the *Borrelia* polypeptide-specific antibodies of the present invention are provided below. Examples of suitable enzyme labels include malate dehydrogenase, *Borrelia* nuclease, delta-5-steroid isomerase, yeast-alcohol dehydrogenase, alpha-glycerol phosphate dehydrogenase, triose phosphate isomerase, peroxidase, alkaline phosphatase, asparaginase, glucose oxidase, beta-galactosidase, ribonuclease, urease, catalase, glucose-6-phosphate dehydrogenase, glucoamylase, and acetylcholine esterase.

Examples of suitable radioisotopic labels include ^3H , ^{111}In , ^{125}I , ^{131}I , ^{32}P , ^{35}S , ^{14}C , ^{51}Cr , ^{57}Co , ^{59}Fe , ^{75}Se , ^{152}Eu , ^{90}Y , ^{67}Cu , ^{217}Cl , ^{211}At , ^{212}Pb , ^{47}Sc , ^{109}Pd , etc. ^{111}In is a preferred isotope where *in vivo* imaging is used since its avoids the problem of dehalogenation of the ^{125}I or ^{131}I -labeled monoclonal antibody by the liver. In addition, this radionucleotide has a more favorable gamma emission energy for imaging. See, e.g., Perkins et al. (1985) *Eur. J. Nucl. Med.* 10:296-301; Carasquillo et al. (1987) *J. Nucl. Med.* 28:281-287. For example, ^{111}In coupled to monoclonal antibodies with 1-(P-isothiocyanatobenzyl)-DPTA has shown little uptake in non-tumors tissues, particularly the liver, and therefore enhances specificity of tumor localization. See, Esteban et al. (1987) *J. Nucl. Med.* 28:861-870.

Examples of suitable non-radioactive isotopic labels include ^{157}Gd , ^{55}Mn , ^{162}Dy , ^{52}Tr , and ^{56}Fe .

Examples of suitable fluorescent labels include an ^{152}Eu label, a fluorescein label, an isothiocyanate label, a rhodamine label, a phycoerythrin label, a phycocyanin label, an allophycocyanin label, an o-phthaldehyde label, and a fluorescamine label.

Examples of suitable toxin labels include, *Pseudomonas* toxin, diphtheria toxin, ricin, and cholera toxin.

Examples of chemiluminescent labels include a luminal label, an isoluminal label, an aromatic acridinium ester label, an imidazole label, an acridinium salt label, an oxalate ester label, a luciferin label, a luciferase label, and an aequorin label.

Examples of nuclear magnetic resonance contrasting agents include heavy metal nuclei such as Gd, Mn, and iron.

Typical techniques for binding the above-described labels to antibodies are provided by

Kennedy et al. (1976) Clin. Chim. Acta 70:1-31, and Schurs et al. (1977) Clin. Chim. Acta 81:1-40. Coupling techniques mentioned in the latter are the glutaraldehyde method, the periodate method, the dimaleimide method, the m-maleimidobenzyl-N-hydroxy-succinimide ester method, all of which methods are incorporated by reference herein.

5 In a related aspect, the invention includes a diagnostic kit for use in screening serum containing antibodies specific against *B. burgdorferi* infection. Such a kit may include an isolated *B. burgdorferi* antigen comprising an epitope which is specifically immunoreactive with at least one anti-*B. burgdorferi* antibody. Such a kit also includes means for detecting the binding of said antibody to the antigen. In specific embodiments, the kit may include a 10 recombinantly produced or chemically synthesized peptide or polypeptide antigen. The peptide or polypeptide antigen may be attached to a solid support.

In a more specific embodiment, the detecting means of the above-described kit includes a solid support to which said peptide or polypeptide antigen is attached. Such a kit may also include a non-attached reporter-labeled anti-human antibody. In this embodiment, binding of the antibody to the *B. burgdorferi* antigen can be detected by binding of the reporter labeled antibody to the anti-*B. burgdorferi* polypeptide antibody.

In a related aspect, the invention includes a method of detecting *B. burgdorferi* infection in a subject. This detection method includes reacting a body fluid, preferably serum, from the subject with an isolated *B. burgdorferi* antigen, and examining the antigen for the presence of bound antibody. In a specific embodiment, the method includes a polypeptide antigen attached to a solid support, and serum is reacted with the support. Subsequently, the support is reacted with a reporter-labeled anti-human antibody. The support is then examined for the presence of reporter-labeled antibody.

The solid surface reagent employed in the above assays and kits is prepared by known 25 techniques for attaching protein material to solid support material, such as polymeric beads, dip sticks, 96-well plates or filter material. These attachment methods generally include non-specific adsorption of the protein to the support or covalent attachment of the protein, typically through a free amine group, to a chemically reactive group on the solid support, such as an activated carboxyl, hydroxyl, or aldehyde group. Alternatively, streptavidin coated plates can be used in 30 conjunction with biotinylated antigen(s).

The polypeptides and antibodies of the present invention, including fragments thereof, 35 may be used to detect *Borrelia* species including *B. burgdorferi* using bio chip and biosensor technology. Bio chip and biosensors of the present invention may comprise the polypeptides of the present invention to detect antibodies, which specifically recognize *Borrelia* species, including *B. burgdorferi*. Bio chip and biosensors of the present invention may also comprise antibodies which specifically recognize the polypeptides of the present invention to detect *Borrelia* species, including *B. burgdorferi* or specific polypeptides of the present invention. Bio chips or biosensors comprising polypeptides or antibodies of the present invention may be used to detect *Borrelia* species, including *B. burgdorferi*, in biological and environmental samples and to

diagnose an animal, including humans, with an *B. burgdorferi* or other *Borrelia* infection. Thus, the present invention includes both bio chips and biosensors comprising polypeptides or antibodies of the present invention and methods of their use.

The bio chips of the present invention may further comprise polypeptide sequences of other pathogens including bacteria, viral, parasitic, and fungal polypeptide sequences, in addition to the polypeptide sequences of the present invention, for use in rapid differential pathogenic detection and diagnosis. The bio chips of the present invention may further comprise antibodies or fragments thereof specific for other pathogens including bacteria, viral, parasitic, and fungal polypeptide sequences, in addition to the antibodies or fragments thereof of the present invention, for use in rapid differential pathogenic detection and diagnosis. The bio chips and biosensors of the present invention may also be used to monitor an *B. burgdorferi* or other *Borrelia* infection and to monitor the genetic changes (amino acid deletions, insertions, substitutions, etc.) in response to drug therapy in the clinic and drug development in the laboratory. The bio chip and biosensors comprising polypeptides or antibodies of the present invention may also be used to simultaneously monitor the expression of a multiplicity of polypeptides, including those of the present invention. The polypeptides used to comprise a bio chip or biosensor of the present invention may be specified in the same manner as for the fragments, i.e., by their N-terminal and C-terminal positions or length in contiguous amino acid residue. Methods and particular uses of the polypeptides and antibodies of the present invention to detect *Borrelia* species, including *B. burgdorferi*, or specific polypeptides using bio chip and biosensor technology include those known in the art, those of the U.S. Patent Nos. and World Patent Nos. listed above for bio chips and biosensors using polynucleotides of the present invention, and those of: U.S. Patent Nos. 5658732, 5135852, 5567301, 5677196, 5690894 and World Patent Nos. WO9729366, WO9612957, each incorporated herein in their entireties.

25.

Treatment:

Agonists and Antagonists - Assays and Molecules

The invention also provides a method of screening compounds to identify those which enhance or block the biological activity of the *B. burgdorferi* polypeptides of the present invention. The present invention further provides where the compounds kill or slow the growth of *B. burgdorferi*. The ability of *B. burgdorferi* antagonists, including *B. burgdorferi* ligands, to prophylactically or therapeutically block antibiotic resistance may be easily tested by the skilled artisan. *See, e.g.*, Straden et al. (1997) J Bacteriol. 179(1):9-16.

An agonist is a compound which increases the natural biological function or which functions in a manner similar to the polypeptides of the present invention, while antagonists decrease or eliminate such functions. Potential antagonists include small organic molecules, peptides, polypeptides, and antibodies that bind to a polypeptide of the invention and thereby inhibit or extinguish its activity.

The antagonists may be employed for instance to inhibit peptidoglycan cross bridge

formation. Antibodies against *B. burgdorferi* may be employed to bind to and inhibit *B. burgdorferi* activity to treat antibiotic resistance. Any of the above antagonists may be employed in a composition with a pharmaceutically acceptable carrier.

5 **Vaccines**

The present invention also provides vaccines comprising one or more polypeptides of the present invention. Heterogeneity in the composition of a vaccine may be provided by combining *B. burgdorferi* polypeptides of the present invention. Multi-component vaccines of this type are desirable because they are likely to be more effective in eliciting protective immune responses 10 against multiple species and strains of the *Borrelia* genus than single polypeptide vaccines. Thus, as discussed in detail below, a multi-component vaccine of the present invention may contain one or more, preferably 2 to about 20, more preferably 2 to about 15, and most preferably 3 to about 8, of the *B. burgdorferi* polypeptides shown in Table 1, or fragments thereof.

Multi-component vaccines are known in the art to elicit antibody production to numerous immunogenic components. Decker, M. and Edwards, K., *J. Infect. Dis.* 174:S270-275 (1996). In addition, a hepatitis B, diphtheria, tetanus, pertussis tetravalent vaccine has recently been demonstrated to elicit protective levels of antibodies in human infants against all four pathogenic agents. Aristegui, J. *et al.*, *Vaccine* 15:7-9 (1997).

The present invention thus also includes multi-component vaccines. These vaccines comprise more than one polypeptide, immunogen or antigen. An example of such a multi-component vaccine would be a vaccine comprising more than one of the *B. burgdorferi* polypeptides shown in Table 1. A second example is a vaccine comprising one or more, for example 2 to 10, of the *B. burgdorferi* polypeptides shown in Table 1 and one or more, for example 2 to 10, additional polypeptides of either borrelial or non-borrelial origin. Thus, a multi-component vaccine which confers protective immunity to both a borrelial infection and infection by another pathogenic agent is also within the scope of the invention.

As indicated above, the vaccines of the present invention are expected to elicit a protective immune response against infections caused by species and strains of *Borrelia* other than *B. burgdorferi* sensu stricto isolate B31 (ATCC Accession No. 35210). Immunizations using 30 decorin-binding protein and OspA derived from one strain of *B. burgdorferi* has been shown to elicit the production of antiserum which confers passive immunity against other strains of *B. burgdorferi*. Cassatt, D. *et al.*, *Protection of Borrelia burgdorferi Infection by Antibodies to Decorin-binding Protein*, in VACCINES97, Cold Spring Harbor Press (1997), pages 191-195. Further, the inventors have found using an *in vitro* assay that antiserum produced in response to 35 *B. burgdorferi* decorin-binding protein will kill several species of *Borrelia*. The amino acid sequences of decorin-binding protein expressed by different strains of *B. burgdorferi* are believed to diverge by as much as 25%. Thus, antisera elicited against decorin-binding proteins confers passive immunity against *Borrelia* expressing proteins having only 75% or less amino acid sequence similarity.

Further within the scope of the invention are whole cell and whole viral vaccines. Such vaccines may be produced recombinantly and involve the expression of one or more of the *B. burgdorferi* polypeptides shown in Table 1. For example, the *B. burgdorferi* polypeptides of the present invention may be either secreted or localized intracellular, on the cell surface, or in the periplasmic space. Further, when a recombinant virus is used, the *B. burgdorferi* polypeptides of the present invention may, for example, be localized in the viral envelope, on the surface of the capsid, or internally within the capsid. Whole cells vaccines which employ cells expressing heterologous proteins are known in the art. *See, e.g.*, Robinson, K. *et al.*, *Nature Biotech.* 15:653-657 (1997); Sirard, J. *et al.*, *Infect. Immun.* 65:2029-2033 (1997); Chabalgoity, J. *et al.*, *Infect. Immun.* 65:2402-2412 (1997). These cells may be administered live or may be killed prior to administration. Chabalgoity, J. *et al.*, *supra*, for example, report the successful use in mice of a live attenuated *Salmonella* vaccine strain which expresses a portion of a platyhelminth fatty acid-binding protein as a fusion protein on its cells surface.

A multi-component vaccine can also be prepared using techniques known in the art by combining one or more *B. burgdorferi* polypeptides of the present invention, or fragments thereof, with additional non-borrelial components (*e.g.*, diphtheria toxin or tetanus toxin, and/or other compounds known to elicit an immune response). Such vaccines are useful for eliciting protective immune responses to both members of the *Borrelia* genus and non-borrelial pathogenic agents.

The vaccines of the present invention also include DNA vaccines. DNA vaccines are currently being developed for a number of infectious diseases. Boyer, J *et al.*, *Nat. Med.* 3:526-532 (1997); reviewed in Spier, R., *Vaccine* 14:1285-1288 (1996). Such DNA vaccines contain a nucleotide sequence encoding one or more *B. burgdorferi* polypeptides of the present invention oriented in a manner that allows for expression of the subject polypeptide. The direct administration of plasmid DNA encoding OspA has been shown to elicit protective immunity in mice against borrelial challenge. Luke, C. *et al.*, *J. Infect. Dis.* 175:91-97 (1997).

The present invention also relates to the administration of a vaccine which is co-administered with a molecule capable of modulating immune responses. Kim, J. *et al.*, *Nature Biotech.* 15:641-646 (1997), for example, report the enhancement of immune responses produced by DNA immunizations when DNA sequences encoding molecules which stimulate the immune response are co-administered. In a similar fashion, the vaccines of the present invention may be co-administered with either nucleic acids encoding immune modulators or the immune modulators themselves. These immune modulators include granulocyte macrophage colony stimulating factor (GM-CSF) and CD86.

The vaccines of the present invention may be used to confer resistance to borrelial infection by either passive or active immunization. When the vaccines of the present invention are used to confer resistance to borrelial infection through active immunization, a vaccine of the present invention is administered to an animal to elicit a protective immune response which either prevents or attenuates a borrelial infection. When the vaccines of the present invention are used to

confer resistance to borrelial infection through passive immunization, the vaccine is provided to a host animal (e.g., human, dog, or mouse), and the antisera elicited by this antisera is recovered and directly provided to a recipient suspected of having an infection caused by a member of the *Borrelia* genus.

5 The ability to label antibodies, or fragments of antibodies, with toxin molecules provides an additional method for treating borrelial infections when passive immunization is conducted. In this embodiment, antibodies, or fragments of antibodies, capable of recognizing the *B. burgdorferi* polypeptides disclosed herein, or fragments thereof, as well as other *Borrelia* proteins, are labeled with toxin molecules prior to their administration to the patient. When such 10 toxin derivatized antibodies bind to *Borrelia* cells, toxin moieties will be localized to these cells and will cause their death.

15 The present invention thus concerns and provides a means for preventing or attenuating a borrelial infection resulting from organisms which have antigens that are recognized and bound by antisera produced in response to the polypeptides of the present invention. As used herein, a vaccine is said to prevent or attenuate a disease if its administration to an animal results either in the total or partial attenuation (i.e., suppression) of a symptom or condition of the disease, or in the total or partial immunity of the animal to the disease.

20 The administration of the vaccine (or the antisera which it elicits) may be for either a "prophylactic" or "therapeutic" purpose. When provided prophylactically, the compound(s) are provided in advance of any symptoms of borrelial infection. The prophylactic administration of the compound(s) serves to prevent or attenuate any subsequent infection. When provided therapeutically, the compound(s) is provided upon or after the detection of symptoms which indicate that an animal may be infected with a member of the *Borrelia* genus. The therapeutic 25 administration of the compound(s) serves to attenuate any actual infection. Thus, the *B. burgdorferi* polypeptides, and fragments thereof, of the present invention may be provided either prior to the onset of infection (so as to prevent or attenuate an anticipated infection) or after the initiation of an actual infection.

30 The polypeptides of the invention, whether encoding a portion of a native protein or a functional derivative thereof, may be administered in pure form or may be coupled to a macromolecular carrier. Examples of such carriers are proteins and carbohydrates. Suitable proteins which may act as macromolecular carrier for enhancing the immunogenicity of the polypeptides of the present invention include keyhole limpet hemocyanin (KLH) tetanus toxoid, pertussis toxin, bovine serum albumin, and ovalbumin. Methods for coupling the polypeptides of the present invention to such macromolecular carriers are disclosed in Harlow *et al.*, *Antibodies: A Laboratory Manual*, 2nd Ed.; Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (1988), the entire disclosure of which is incorporated by reference herein.

35 A composition is said to be "pharmacologically acceptable" if its administration can be tolerated by a recipient animal and is otherwise suitable for administration to that animal. Such an agent is said to be administered in a "therapeutically effective amount" if the amount administered

is physiologically significant. An agent is physiologically significant if its presence results in a detectable change in the physiology of a recipient patient.

While in all instances the vaccine of the present invention is administered as a pharmacologically acceptable compound, one skilled in the art would recognize that the composition of a pharmacologically acceptable compound varies with the animal to which it is administered. For example, a vaccine intended for human use will generally not be co-administered with Freund's adjuvant. Further, the level of purity of the *B. burgdorferi* polypeptides of the present invention will normally be higher when administered to a human than when administered to a non-human animal.

As would be understood by one of ordinary skill in the art, when the vaccine of the present invention is provided to an animal, it may be in a composition which may contain salts, buffers, adjuvants, or other substances which are desirable for improving the efficacy of the composition. Adjuvants are substances that can be used to specifically augment a specific immune response. These substances generally perform two functions: (1) they protect the antigen(s) from being rapidly catabolized after administration and (2) they nonspecifically stimulate immune responses.

Normally, the adjuvant and the composition are mixed prior to presentation to the immune system, or presented separately, but into the same site of the animal being immunized. Adjuvants can be loosely divided into several groups based upon their composition. These groups include oil adjuvants (for example, Freund's complete and incomplete), mineral salts (for example, $AIK(SO_4)_2$, $AlNa(SO_4)_2$, $AlNH_4(SO_4)_2$, silica, kaolin, and carbon), polynucleotides (for example, poly IC and poly AU acids), and certain natural substances (for example, wax D from *Mycobacterium tuberculosis*, as well as substances found in *Corynebacterium parvum*, or *Bordetella pertussis*, and members of the genus *Brucella*. Other substances useful as adjuvants are the saponins such as, for example, Quil A. (Superfos A/S, Denmark). Preferred adjuvants for use in the present invention include aluminum salts, such as $AIK(SO_4)_2$, $AlNa(SO_4)_2$, and $AlNH_4(SO_4)_2$. Examples of materials suitable for use in vaccine compositions are provided in *Remington's Pharmaceutical Sciences* (Osol, A, Ed, Mack Publishing Co, Easton, PA, pp. 1324-1341 (1980), which reference is incorporated herein by reference).

The therapeutic compositions of the present invention can be administered parenterally by injection, rapid infusion, nasopharyngeal absorption (intranasopharangeally), dermoabsorption, or orally. The compositions may alternatively be administered intramuscularly, or intravenously. Compositions for parenteral administration include sterile aqueous or non-aqueous solutions, suspensions, and emulsions. Examples of non-aqueous solvents are propylene glycol, polyethylene glycol, vegetable oils such as olive oil, and injectable organic esters such as ethyl oleate. Carriers or occlusive dressings can be used to increase skin permeability and enhance antigen absorption. Liquid dosage forms for oral administration may generally comprise a liposome solution containing the liquid dosage form. Suitable forms for suspending liposomes include emulsions, suspensions, solutions, syrups, and elixirs containing inert diluents

commonly used in the art, such as purified water. Besides the inert diluents, such compositions can also include adjuvants, wetting agents, emulsifying and suspending agents, or sweetening, flavoring, or perfuming agents.

Therapeutic compositions of the present invention can also be administered in encapsulated form. For example, intranasal immunization of mice against *Bordetella pertussis* infection using vaccines encapsulated in biodegradable microsphere composed of poly(DL-lactide-co-glycolide) has been shown to stimulate protective immune responses. Shahin, R. *et al.*, *Infect. Immun.* 63:1195-1200 (1995). Similarly, orally administered encapsulated *Salmonella typhimurium* antigens have also been shown to elicit protective immunity in mice. Allaoui-Attarki, K. *et al.*, *Infect. Immun.* 65:853-857 (1997). Encapsulated vaccines of the present invention can be administered by a variety of routes including those involving contacting the vaccine with mucous membranes (e.g., intranasally, intracolonically, intraduodenally).

Many different techniques exist for the timing of the immunizations when a multiple administration regimen is utilized. It is possible to use the compositions of the invention more than once to increase the levels and diversities of expression of the immunoglobulin repertoire expressed by the immunized animal. Typically, if multiple immunizations are given, they will be given one to two months apart.

According to the present invention, an "effective amount" of a therapeutic composition is one which is sufficient to achieve a desired biological effect. Generally, the dosage needed to provide an effective amount of the composition will vary depending upon such factors as the animal's or human's age, condition, sex, and extent of disease, if any, and other variables which can be adjusted by one of ordinary skill in the art.

The antigenic preparations of the invention can be administered by either single or multiple dosages of an effective amount. Effective amounts of the compositions of the invention can vary from 0.01-1,000 µg/ml per dose, more preferably 0.1-500 µg/ml per dose, and most preferably 10-300 µg/ml per dose.

Having now generally described the invention, the same will be more readily understood through reference to the following example which is provided by way of illustration, and is not intended to be limiting of the present invention, unless specified.

Examples

1. Preparation of PCR Primers and Amplification of DNA

Various fragments of the *Borrelia burgdorferi* genome, such as those of Table 1, can be used, in accordance with the present invention, to prepare PCR primers for a variety of uses. The PCR primers are preferably at least 15 bases, and more preferably at least 18 bases in length. When selecting a primer sequence, it is preferred that the primer pairs have approximately the same G/C ratio, so that melting temperatures are approximately the same. The PCR primers and

amplified DNA of this Example find use in the Examples that follow.

2. Isolation of a Selected DNA Clone From *B. burgdorferi*

Three approaches are used to isolate a *B. burgdorferi* clone comprising a polynucleotide of the present invention from any *B. burgdorferi* genomic DNA library. The *B. burgdorferi* strain B31PU has been deposited as a convenient source for obtaining a *B. burgdorferi* strain although a wide variety of strains *B. burgdorferi* strains can be used which are known in the art.

B. burgdorferi genomic DNA is prepared using the following method. A 20ml overnight bacterial culture grown in a rich medium (e.g., Trypticase Soy Broth, Brain Heart Infusion broth or Super broth), pelleted, ished two times with TES (30mM Tris-pH 8.0, 25mM EDTA, 50mM NaCl), and resuspended in 5ml high salt TES (2.5M NaCl). Lysostaphin is added to final concentration of approx 50ug/ml and the mixture is rotated slowly 1 hour at 37C to make protoplast cells. The solution is then placed in incubator (or place in a shaking water bath) and warmed to 55C. Five hundred micro liter of 20% sarcosyl in TES (final concentration 2%) is then added to lyse the cells. Next, guanidine HCl is added to a final concentration of 7M (3.69g in 5.5 ml). The mixture is swirled slowly at 55C for 60-90 min (solution should clear). A CsCl gradient is then set up in SW41 ultra-clear tubes using 2.0ml 5.7M CsCl and overlaying with 2.85M CsCl. The gradient is carefully overlayed with the DNA-containing GuHCl solution. The gradient is spun at 30,000 rpm, 20C for 24 hr and the lower DNA band is collected. The volume is increased to 5 ml with TE buffer. The DNA is then treated with protease K (10 ug/ml) overnight at 37 C, and precipitated with ethanol. The precipitated DNA is resuspended in a desired buffer.

In the first method, a plasmid is directly isolated by screening a plasmid *B. burgdorferi* genomic DNA library using a polynucleotide probe corresponding to a polynucleotide of the present invention. Particularly, a specific polynucleotide with 30-40 nucleotides is synthesized using an Applied Biosystems DNA synthesizer according to the sequence reported. The oligonucleotide is labeled, for instance, with ^{32}P - γ -ATP using T4 polynucleotide kinase and purified according to routine methods. (See, e.g., Maniatis et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Press, Cold Spring, NY (1982).) The library is transformed into a suitable host, as indicated above (such as XL-1 Blue (Stratagene)) using techniques known to those of skill in the art. See, e.g., Sambrook et al. MOLECULAR CLONING: A LABORATORY MANUAL (Cold Spring Harbor, N.Y. 2nd ed. 1989); Ausubel et al., CURRENT PROTOCOLS IN MOLECULAR BIOLOGY (John Wiley and Sons, N.Y. 1989). The transformants are plated on 1.5% agar plates (containing the appropriate selection agent, e.g., ampicillin) to a density of about 150 transformants (colonies) per plate. These plates are screened using Nylon membranes according to routine methods for bacterial colony screening. See, e.g., Sambrook et al. MOLECULAR CLONING: A LABORATORY MANUAL (Cold Spring Harbor, N.Y. 2nd ed. 1989); Ausubel et al., CURRENT PROTOCOLS IN

MOLECULAR BIOLOGY (John Wiley and Sons, N.Y. 1989) or other techniques known to those of skill in the art.

Alternatively, two primers of 15-25 nucleotides derived from the 5' and 3' ends of a polynucleotide of Table 1 are synthesized and used to amplify the desired DNA by PCR using a *B. burgdorferi* genomic DNA prep as a template. PCR is carried out under routine conditions, for instance, in 25 μ l of reaction mixture with 0.5 ug of the above DNA template. A convenient reaction mixture is 1.5-5 mM MgCl₂, 0.01% (w/v) gelatin, 20 μ M each of dATP, dCTP, dGTP, dTTP, 25 pmol of each primer and 0.25 Unit of Taq polymerase. Thirty five cycles of PCR (denaturation at 94°C for 1 min; annealing at 55°C for 1 min; elongation at 72°C for 1 min) are performed with a Perkin-Elmer Cetus automated thermal cycler. The amplified product is analyzed by agarose gel electrophoresis and the DNA band with expected molecular weight is excised and purified. The PCR product is verified to be the selected sequence by subcloning and sequencing the DNA product.

Finally, overlapping oligos of the DNA sequences of Table 1 can be chemically synthesized and used to generate a nucleotide sequence of desired length using PCR methods known in the art.

3(a). *Expression and Purification Borrelia polypeptides in E. coli*

The bacterial expression vector pQE60 is used for bacterial expression of some of the polypeptide fragments of the present invention. (QIAGEN, Inc., 9259 Eton Avenue, Chatsworth, CA, 91311). pQE60 encodes ampicillin antibiotic resistance ("Amp^r") and contains a bacterial origin of replication ("ori"), an IPTG inducible promoter, a ribosome binding site ("RBS"), six codons encoding histidine residues that allow affinity purification using nickel-nitrilo-tri-acetic acid ("Ni-NTA") affinity resin (QIAGEN, Inc., *supra*) and suitable single restriction enzyme cleavage sites. These elements are arranged such that an inserted DNA fragment encoding a polypeptide expresses that polypeptide with the six His residues (i.e., a "6 X His tag") covalently linked to the carboxyl terminus of that polypeptide.

The DNA sequence encoding the desired portion of a *B. burgdorferi* protein of the present invention is amplified from *B. burgdorferi* genomic DNA using PCR oligonucleotide primers which anneal to the 5' and 3' sequences coding for the portions of the *B. burgdorferi* polynucleotide shown in Table 1. Additional nucleotides containing restriction sites to facilitate cloning in the pQE60 vector are added to the 5' and 3' sequences, respectively.

For cloning the mature protein, the 5' primer has a sequence containing an appropriate restriction site followed by nucleotides of the amino terminal coding sequence of the desired *B. burgdorferi* polynucleotide sequence in Table 1. One of ordinary skill in the art would appreciate that the point in the protein coding sequence where the 5' and 3' primers begin may be varied to amplify a DNA segment encoding any desired portion of the complete protein shorter or longer than the mature form. The 3' primer has a sequence containing an appropriate restriction site

followed by nucleotides complementary to the 3' end of the polypeptide coding sequence of Table 1, excluding a stop codon, with the coding sequence aligned with the restriction site so as to maintain its reading frame with that of the six His codons in the pQE60 vector.

The amplified *B. burgdorferi* DNA fragment and the vector pQE60 are digested with restriction enzymes which recognize the sites in the primers and the digested DNAs are then ligated together. The *B. burgdorferi* DNA is inserted into the restricted pQE60 vector in a manner which places the *B. burgdorferi* protein coding region downstream from the IPTG-inducible promoter and in-frame with an initiating AUG and the six histidine codons.

The ligation mixture is transformed into competent *E. coli* cells using standard procedures such as those described by Sambrook et al., *supra*. *E. coli* strain M15/rep4, containing multiple copies of the plasmid pREP4, which expresses the lac repressor and confers kanamycin resistance ("Kanr"), is used in carrying out the illustrative example described herein. This strain, which is only one of many that are suitable for expressing a *B. burgdorferi* polypeptide, is available commercially (QIAGEN, Inc., *supra*). Transformants are identified by their ability to grow on LB agar plates in the presence of ampicillin and kanamycin. Plasmid DNA is isolated from resistant colonies and the identity of the cloned DNA confirmed by restriction analysis, PCR and DNA sequencing.

Clones containing the desired constructs are grown overnight ("O/N") in liquid culture in LB media supplemented with both ampicillin (100 µg/ml) and kanamycin (25 µg/ml). The O/N culture is used to inoculate a large culture, at a dilution of approximately 1:25 to 1:250. The cells are grown to an optical density at 600 nm ("OD600") of between 0.4 and 0.6. Isopropyl-β-D-thiogalactopyranoside ("IPTG") is then added to a final concentration of 1 mM to induce transcription from the lac repressor sensitive promoter, by inactivating the lacI repressor. Cells subsequently are incubated further for 3 to 4 hours. Cells then are harvested by centrifugation.

The cells are then stirred for 3-4 hours at 4°C in 6M guanidine-HCl, pH 8. The cell debris is removed by centrifugation, and the supernatant containing the *B. burgdorferi* polypeptide is loaded onto a nickel-nitrolo-tri-acetic acid ("Ni-NTA") affinity resin column (QIAGEN, Inc., *supra*). Proteins with a 6 x His tag bind to the Ni-NTA resin with high affinity are purified in a simple one-step procedure (for details see: The QIAexpressionist, 1995, QIAGEN, Inc., *supra*). Briefly the supernatant is loaded onto the column in 6 M guanidine-HCl, pH 8, the column is first washed with 10 volumes of 6 M guanidine-HCl, pH 8, then washed with 10 volumes of 6 M guanidine-HCl pH 6, and finally the *B. burgdorferi* polypeptide is eluted with 6 M guanidine-HCl, pH 5.

The purified protein is then renatured by dialyzing it against phosphate-buffered saline (PBS) or 50 mM Na-acetate, pH 6 buffer plus 200 mM NaCl. Alternatively, the protein could be successfully refolded while immobilized on the Ni-NTA column. The recommended conditions are as follows: renature using a linear 6M-1M urea gradient in 500 mM NaCl, 20% glycerol, 20 mM Tris/HCl pH 7.4, containing protease inhibitors. The renaturation should be performed over

a period of 1.5 hours or more. After renaturation the proteins can be eluted by the addition of 250 mM immidazole. Immidazole is removed by a final dialyzing step against PBS or 50 mM sodium acetate pH 6 buffer plus 200 mM NaCl. The purified protein is stored at 4° C or frozen at -80° C.

The polypeptide of the present invention are also prepared using a non-denaturing protein

5 purification method. For these polypeptides, the cell pellet from each liter of culture is resuspended in 25 mls of Lysis Buffer A at 4°C (Lysis Buffer A = 50 mM Na-phosphate, 300 mM NaCl, 10 mM 2-mercaptoethanol, 10% Glycerol, pH 7.5 with 1 tablet of Complete EDTA-free protease inhibitor cocktail (Boehringer Mannheim #1873580) per 50 ml of buffer).

10 Absorbance at 550 nm is approximately 10-20 O.D./ml. The suspension is then put through three freeze/thaw cycles from -70°C (using a ethanol-dry ice bath) up to room temperature. The cells are lysed via sonication in short 10 sec bursts over 3 minutes at approximately 80W while kept on ice. The sonicated sample is then centrifuged at 15,000 RPM for 30 minutes at 4°C. The supernatant is passed through a column containing 1.0 ml of CL-4B resin to pre-clear the sample of any proteins that may bind to agarose non-specifically, and the flow-through fraction is collected.

15 The pre-cleared flow-through is applied to a nickel-nitrilo-tri-acetic acid ("Ni-NTA") affinity resin column (Quiagen, Inc., *supra*). Proteins with a 6 X His tag bind to the Ni-NTA resin with high affinity and can be purified in a simple one-step procedure. Briefly, the supernatant is loaded onto the column in Lysis Buffer A at 4°C, the column is first washed with 20 10 volumes of Lysis Buffer A until the A280 of the eluate returns to the baseline. Then, the column is washed with 5 volumes of 40 mM Imidazole (92% Lysis Buffer A / 8% Buffer B) (Buffer B = 50 mM Na-Phosphate, 300 mM NaCl, 10% Glycerol, 10 mM 2-mercaptoethanol, 500 mM Imidazole, pH of the final buffer should be 7.5). The protein is eluted off of the column with a series of increasing Imidazole solutions made by adjusting the ratios of Lysis Buffer A to 25 Buffer B. Three different concentrations are used: 3 volumes of 75 mM Imidazole, 3 volumes of 150 mM Imidazole, 5 volumes of 500 mM Imidazole. The fractions containing the purified protein are analyzed using 8 %, 10 % or 14% SDS-PAGE depending on the protein size. The purified protein is then dialyzed 2X against phosphate-buffered saline (PBS) in order to place it into an easily workable buffer. The purified protein is stored at 4° C or frozen at -80°.

30 The following alternative method may be used to purify *B. burgdorferi* expressed in *E. coli* when it is present in the form of inclusion bodies. Unless otherwise specified, all of the following steps are conducted at 4-10°C.

Upon completion of the production phase of the *E. coli* fermentation, the cell culture is cooled to 4-10°C and the cells are harvested by continuous centrifugation at 15,000 rpm (Heraeus 35 Sepatech). On the basis of the expected yield of protein per unit weight of cell paste and the amount of purified protein required, an appropriate amount of cell paste, by weight, is suspended in a buffer solution containing 100 mM Tris, 50 mM EDTA, pH 7.4. The cells are dispersed to a homogeneous suspension using a high shear mixer.

The cells are then lysed by passing the solution through a microfluidizer (Microfluidics, Corp. or APV Gaulin, Inc.) twice at 4000-6000 psi. The homogenate is then mixed with NaCl solution to a final concentration of 0.5 M NaCl, followed by centrifugation at 7000 x g for 15 min. The resultant pellet is washed again using 0.5M NaCl, 100 mM Tris, 50 mM EDTA, pH 5 7.4.

The resulting washed inclusion bodies are solubilized with 1.5 M guanidine hydrochloride (GuHCl) for 2-4 hours. After 7000 x g centrifugation for 15 min., the pellet is discarded and the *B. burgdorferi* polypeptide-containing supernatant is incubated at 4°C overnight to allow further GuHCl extraction.

Following high speed centrifugation (30,000 x g) to remove insoluble particles, the GuHCl solubilized protein is refolded by quickly mixing the GuHCl extract with 20 volumes of buffer containing 50 mM sodium, pH 4.5, 150 mM NaCl, 2 mM EDTA by vigorous stirring. The refolded diluted protein solution is kept at 4°C without mixing for 12 hours prior to further purification steps.

To clarify the refolded *B. burgdorferi* polypeptide solution, a previously prepared tangential filtration unit equipped with 0.16 µm membrane filter with appropriate surface area (e.g., Filtron), equilibrated with 40 mM sodium acetate, pH 6.0 is employed. The filtered sample is loaded onto a cation exchange resin (e.g., Poros HS-50, Perseptive Biosystems). The column is washed with 40 mM sodium acetate, pH 6.0 and eluted with 250 mM, 500 mM, 1000 mM, and 1500 mM NaCl in the same buffer, in a stepwise manner. The absorbance at 280 nm of the effluent is continuously monitored. Fractions are collected and further analyzed by SDS-PAGE.

Fractions containing the *B. burgdorferi* polypeptide are then pooled and mixed with 4 volumes of water. The diluted sample is then loaded onto a previously prepared set of tandem columns of strong anion (Poros HQ-50, Perseptive Biosystems) and weak anion (Poros CM-20, Perseptive Biosystems) exchange resins. The columns are equilibrated with 40 mM sodium acetate, pH 6.0. Both columns are washed with 40 mM sodium acetate, pH 6.0, 200 mM NaCl. The CM-20 column is then eluted using a 10 column volume linear gradient ranging from 0.2 M NaCl, 50 mM sodium acetate, pH 6.0 to 1.0 M NaCl, 50 mM sodium acetate, pH 6.5. Fractions are collected under constant A_{280} monitoring of the effluent. Fractions containing the *B. burgdorferi* polypeptide (determined, for instance, by 16% SDS-PAGE) are then pooled.

The resultant *B. burgdorferi* polypeptide exhibits greater than 95% purity after the above refolding and purification steps. No major contaminant bands are observed from Commassie blue stained 16% SDS-PAGE gel when 5 µg of purified protein is loaded. The purified protein is also tested for endotoxin/LPS contamination, and typically the LPS content is less than 0.1 ng/ml according to LAL assays.

coli

The vector pQE10 is alternatively used to clone and express some of the polypeptides of the present invention for use in the soft tissue and systemic infection models discussed below. The difference being such that an inserted DNA fragment encoding a polypeptide expresses that polypeptide with the six His residues (i.e., a "6 X His tag") covalently linked to the amino terminus of that polypeptide. The bacterial expression vector pQE10 (QIAGEN, Inc., 9259 Eton Avenue, Chatsworth, CA, 91311) was used in this example. The components of the pQE10 plasmid are arranged such that the inserted DNA sequence encoding a polypeptide of the present invention expresses the polypeptide with the six His residues (i.e., a "6 X His tag") covalently linked to the amino terminus.

The DNA sequences encoding the desired portions of a polypeptide of Table 1 were amplified using PCR oligonucleotide primers from genomic *B. burgdorferi* DNA. The PCR primers anneal to the nucleotide sequences encoding the desired amino acid sequence of a polypeptide of the present invention. Additional nucleotides containing restriction sites to facilitate cloning in the pQE10 vector were added to the 5' and 3' primer sequences, respectively.

For cloning a polypeptide of the present invention, the 5' and 3' primers were selected to amplify their respective nucleotide coding sequences. One of ordinary skill in the art would appreciate that the point in the protein coding sequence where the 5' and 3' primers begins may be varied to amplify a DNA segment encoding any desired portion of a polypeptide of the present invention. The 5' primer was designed so the coding sequence of the 6 X His tag is aligned with the restriction site so as to maintain its reading frame with that of *B. burgdorferi* polypeptide. The 3' was designed to include an stop codon. The amplified DNA fragment was then cloned, and the protein expressed, as described above for the pQE60 plasmid.

The DNA sequences of Table 1 encoding amino acid sequences may also be cloned and expressed as fusion proteins by a protocol similar to that described directly above, wherein the pET-32b(+) vector (Novagen, 601 Science Drive, Madison, WI 53711) is preferentially used in place of pQE10.

The above methods are not limited to the polypeptide fragments actually produced. The above method, like the methods below, can be used to produce either full length polypeptides or desired fragments thereof.

*3(c). Alternative Expression and Purification of *Borrelia* polypeptides in *E. coli*.*

The bacterial expression vector pQE60 is used for bacterial expression in this example (QIAGEN, Inc., 9259 Eton Avenue, Chatsworth, CA, 91311). However, in this example, the polypeptide coding sequence is inserted such that translation of the six His codons is prevented and, therefore, the polypeptide is produced with no 6 X His tag.

The DNA sequence encoding the desired portion of the *B. burgdorferi* amino acid sequence is amplified from an *B. burgdorferi* genomic DNA prep the deposited DNA clones

using PCR oligonucleotide primers which anneal to the 5' and 3' nucleotide sequences corresponding to the desired portion of the *B. burgdorferi* polypeptides. Additional nucleotides containing restriction sites to facilitate cloning in the pQE60 vector are added to the 5' and 3' primer sequences.

5. For cloning a *B. burgdorferi* polypeptides of the present invention, 5' and 3' primers are selected to amplify their respective nucleotide coding sequences. One of ordinary skill in the art would appreciate that the point in the protein coding sequence where the 5' and 3' primers begin may be varied to amplify a DNA segment encoding any desired portion of a polypeptide of the present invention. The 3' and 5' primers contain appropriate restriction sites followed by 10 nucleotides complementary to the 5' and 3' ends of the coding sequence respectively. The 3' primer is additionally designed to include an in-frame stop codon.

The amplified *B. burgdorferi* DNA fragments and the vector pQE60 are digested with restriction enzymes recognizing the sites in the primers and the digested DNAs are then ligated together. Insertion of the *B. burgdorferi* DNA into the restricted pQE60 vector places the *B. burgdorferi* protein coding region including its associated stop codon downstream from the IPTG-inducible promoter and in-frame with an initiating AUG. The associated stop codon prevents translation of the six histidine codons downstream of the insertion point.

20 The ligation mixture is transformed into competent *E. coli* cells using standard procedures such as those described by Sambrook et al. *E. coli* strain M15/rep4, containing multiple copies of the plasmid pREP4, which expresses the lac repressor and confers kanamycin resistance ("Kanr"), is used in carrying out the illustrative example described herein. This strain, which is only one of many that are suitable for expressing *B. burgdorferi* polypeptide, is available commercially (QIAGEN, Inc., *supra*). Transformants are identified by their ability to grow on LB plates in the presence of ampicillin and kanamycin. Plasmid DNA is isolated from resistant 25 colonies and the identity of the cloned DNA confirmed by restriction analysis, PCR and DNA sequencing.

30 Clones containing the desired constructs are grown overnight ("O/N") in liquid culture in LB media supplemented with both ampicillin (100 μ g/ml) and kanamycin (25 μ g/ml). The O/N culture is used to inoculate a large culture, at a dilution of approximately 1:25 to 1:250. The cells are grown to an optical density at 600 nm ("OD600") of between 0.4 and 0.6. isopropyl- β -D-thiogalactopyranoside ("IPTG") is then added to a final concentration of 1 mM to induce transcription from the lac repressor sensitive promoter, by inactivating the lacI repressor. Cells subsequently are incubated further for 3 to 4 hours. Cells then are harvested by centrifugation.

35 To purify the *B. burgdorferi* polypeptide, the cells are then stirred for 3-4 hours at 4°C in 6M guanidine-HCl, pH 8. The cell debris is removed by centrifugation, and the supernatant containing the *B. burgdorferi* polypeptide is dialyzed against 50 mM Na-acetate buffer pH 6, supplemented with 200 mM NaCl. Alternatively, the protein can be successfully refolded by dialyzing it against 500 mM NaCl, 20% glycerol, 25 mM Tris/HCl pH 7.4, containing protease

inhibitors. After renaturation the protein can be purified by ion exchange, hydrophobic interaction and size exclusion chromatography. Alternatively, an affinity chromatography step such as an antibody column can be used to obtain pure *B. burgdorferi* polypeptide. The purified protein is stored at 4°C or frozen at -80°C.

5 The following alternative method may be used to purify *B. burgdorferi* polypeptides expressed in *E. coli* when it is present in the form of inclusion bodies. Unless otherwise specified, all of the following steps are conducted at 4-10°C.

Upon completion of the production phase of the *E. coli* fermentation, the cell culture is cooled to 4-10°C and the cells are harvested by continuous centrifugation at 15,000 rpm (Heraeus 10 Sepatech). On the basis of the expected yield of protein per unit weight of cell paste and the amount of purified protein required, an appropriate amount of cell paste, by weight, is suspended in a buffer solution containing 100 mM Tris, 50 mM EDTA, pH 7.4. The cells are dispersed to a homogeneous suspension using a high shear mixer.

The cells are then lysed by passing the solution through a microfluidizer (Microfluidics, Corp. or APV Gaulin, Inc.) twice at 4000-6000 psi. The homogenate is then mixed with NaCl solution to a final concentration of 0.5 M NaCl, followed by centrifugation at 7000 x g for 15 min. The resultant pellet is washed again using 0.5M NaCl, 100 mM Tris, 50 mM EDTA, pH 7.4.

15 The resulting washed inclusion bodies are solubilized with 1.5 M guanidine hydrochloride (GuHCl) for 2-4 hours. After 7000 x g centrifugation for 15 min., the pellet is discarded and the *B. burgdorferi* polypeptide-containing supernatant is incubated at 4°C overnight to allow further GuHCl extraction.

20 Following high speed centrifugation (30,000 x g) to remove insoluble particles, the GuHCl solubilized protein is refolded by quickly mixing the GuHCl extract with 20 volumes of buffer containing 50 mM sodium, pH 4.5, 150 mM NaCl, 2 mM EDTA by vigorous stirring. The refolded diluted protein solution is kept at 4°C without mixing for 12 hours prior to further purification steps.

25 To clarify the refolded *B. burgdorferi* polypeptide solution, a previously prepared tangential filtration unit equipped with 0.16 µm membrane filter with appropriate surface area (e.g., Filtron), equilibrated with 40 mM sodium acetate, pH 6.0 is employed. The filtered sample is loaded onto a cation exchange resin (e.g., Poros HS-50, Perseptive Biosystems). The column is washed with 40 mM sodium acetate, pH 6.0 and eluted with 250 mM, 500 mM, 1000 mM, and 1500 mM NaCl in the same buffer, in a stepwise manner. The absorbance at 280 nm of the effluent is continuously monitored. Fractions are collected and further analyzed by SDS-PAGE.

30 Fractions containing the *B. burgdorferi* polypeptide are then pooled and mixed with 4 volumes of water. The diluted sample is then loaded onto a previously prepared set of tandem columns of strong anion (Poros HQ-50, Perseptive Biosystems) and weak anion (Poros CM-20,

Perseptive Biosystems) exchange resins. The columns are equilibrated with 40 mM sodium acetate, pH 6.0. Both columns are washed with 40 mM sodium acetate, pH 6.0, 200 mM NaCl. The CM-20 column is then eluted using a 10 column volume linear gradient ranging from 0.2 M NaCl, 50 mM sodium acetate, pH 6.0 to 1.0 M NaCl, 50 mM sodium acetate, pH 6.5. Fractions are collected under constant A_{280} monitoring of the effluent. Fractions containing the *B. burgdorferi* polypeptide (determined, for instance, by 16% SDS-PAGE) are then pooled.

The resultant *B. burgdorferi* polypeptide exhibits greater than 95% purity after the above refolding and purification steps. No major contaminant bands are observed from Commassie blue stained 16% SDS-PAGE gel when 5 μ g of purified protein is loaded. The purified protein is also tested for endotoxin/LPS contamination, and typically the LPS content is less than 0.1 ng/ml according to LAL assays.

3(d). *Cloning and Expression of B. burgdorferi in Other Bacteria*

B. burgdorferi polypeptides can also be produced in: *B. burgdorferi* using the methods of S. Skinner et al., (1988) Mol. Microbiol. 2:289-297 or J. I. Moreno (1996) Protein Expr. Purif. 8(3):332-340; *Lactobacillus* using the methods of C. Rush et al., 1997 Appl. Microbiol. Biotechnol. 47(5):537-542; or in *Bacillus subtilis* using the methods Chang et al., U.S. Patent No. 4,952,508.

4. *Cloning and Expression in COS Cells*

A *B. burgdorferi* expression plasmid is made by cloning a portion of the DNA encoding a *B. burgdorferi* polypeptide into the expression vector pDNAI/Amp or pDNAIII (which can be obtained from Invitrogen, Inc.). The expression vector pDNAI/amp contains: (1) an *E. coli* origin of replication effective for propagation in *E. coli* and other prokaryotic cells; (2) an ampicillin resistance gene for selection of plasmid-containing prokaryotic cells; (3) an SV40 origin of replication for propagation in eukaryotic cells; (4) a CMV promoter, a polylinker, an SV40 intron; (5) several codons encoding a hemagglutinin fragment (i.e., an "HA" tag to facilitate purification) followed by a termination codon and polyadenylation signal arranged so that a DNA can be conveniently placed under expression control of the CMV promoter and operably linked to the SV40 intron and the polyadenylation signal by means of restriction sites in the polylinker. The HA tag corresponds to an epitope derived from the influenza hemagglutinin protein described by Wilson et al. 1984 Cell 37:767. The fusion of the HA tag to the target protein allows easy detection and recovery of the recombinant protein with an antibody that recognizes the HA epitope. pDNAIII contains, in addition, the selectable neomycin marker.

A DNA fragment encoding a *B. burgdorferi* polypeptide is cloned into the polylinker region of the vector so that recombinant protein expression is directed by the CMV promoter. The plasmid construction strategy is as follows. The DNA from a *B. burgdorferi* genomic DNA prep is amplified using primers that contain convenient restriction sites, much as described above for

construction of vectors for expression of *B. burgdorferi* in *E. coli*. The 5' primer contains a Kozak sequence, an AUG start codon, and nucleotides of the 5' coding region of the *B. burgdorferi* polypeptide. The 3' primer, contains nucleotides complementary to the 3' coding sequence of the *B. burgdorferi* DNA, a stop codon, and a convenient restriction site.

5 The PCR amplified DNA fragment and the vector, pDNA1/Amp, are digested with appropriate restriction enzymes and then ligated. The ligation mixture is transformed into an appropriate *E. coli* strain such as SURE™ (Stratagene Cloning Systems, La Jolla, CA 92037), and the transformed culture is plated on ampicillin media plates which then are incubated to allow growth of ampicillin resistant colonies. Plasmid DNA is isolated from resistant colonies and 10 examined by restriction analysis or other means for the presence of the fragment encoding the *B. burgdorferi* polypeptide

For expression of a recombinant *B. burgdorferi* polypeptide, COS cells are transfected with an expression vector, as described above, using DEAE-dextran, as described, for instance, by Sambrook et al. (*supra*). Cells are incubated under conditions for expression of *B. burgdorferi* by the vector.

Expression of the *B. burgdorferi*-HA fusion protein is detected by radiolabeling and immunoprecipitation, using methods described in, for example Harlow et al., *supra*. To this end, two days after transfection, the cells are labeled by incubation in media containing ³⁵S-cysteine for 8 hours. The cells and the media are collected, and the cells are washed and lysed with detergent-containing RIPA buffer: 150 mM NaCl, 1% NP-40, 0.1% SDS, 1% NP-40, 0.5% DOC, 50 mM TRIS, pH 7.5, as described by Wilson et al. (*supra*). Proteins are precipitated from the cell lysate and from the culture media using an HA-specific monoclonal antibody. The precipitated proteins then are analyzed by SDS-PAGE and autoradiography. An expression product of the expected size is seen in the cell lysate, which is not seen in negative controls.

25

5. Cloning and Expression in CHO Cells

The vector pC4 is used for the expression of *B. burgdorferi* polypeptide in this example. Plasmid pC4 is a derivative of the plasmid pSV2-dhfr (ATCC Accession No. 37146). The plasmid contains the mouse DHFR gene under control of the SV40 early promoter. Chinese hamster ovary cells or other cells lacking dihydrofolate activity that are transfected with these plasmids can be selected by growing the cells in a selective medium (alpha minus MEM, Life Technologies) supplemented with the chemotherapeutic agent methotrexate. The amplification of the DHFR genes in cells resistant to methotrexate (MTX) has been well documented. See, e.g., Alt et al., 1978, J. Biol. Chem. 253:1357-1370; Hamlin et al., 1990, Biochem. et Biophys. Acta, 1097:107-143; Page et al., 1991, Biotechnology 9:64-68. Cells grown in increasing concentrations of MTX develop resistance to the drug by overproducing the target enzyme, DHFR, as a result of amplification of the DHFR gene. If a second gene is linked to the DHFR gene, it is usually co-amplified and over-expressed. It is known in the art that this approach may

be used to develop cell lines carrying more than 1,000 copies of the amplified gene(s). Subsequently, when the methotrexate is withdrawn, cell lines are obtained which contain the amplified gene integrated into one or more chromosome(s) of the host cell.

Plasmid pC4 contains the strong promoter of the long terminal repeat (LTR) of the Rouse Sarcoma Virus, for expressing a polypeptide of interest, Cullen, et al. (1985) Mol. Cell. Biol. 5:438-447; plus a fragment isolated from the enhancer of the immediate early gene of human cytomegalovirus (CMV), Boshart, et al., 1985, Cell 41:521-530. Downstream of the promoter are the following single restriction enzyme cleavage sites that allow the integration of the genes: *Bam* HI, *Xba* I, and *Asp* 718. Behind these cloning sites the plasmid contains the 3' intron and polyadenylation site of the rat preproinsulin gene. Other high efficiency promoters can also be used for the expression, e.g., the human β -actin promoter, the SV40 early or late promoters or the long terminal repeats from other retroviruses, e.g., HTV and HTLV. Clontech's Tet-Off and Tet-On gene expression systems and similar systems can be used to express the *B. burgdorferi* polypeptide in a regulated way in mammalian cells (Gossen et al., 1992, Proc. Natl. Acad. Sci. USA 89:5547-5551. For the polyadenylation of the mRNA other signals, e.g., from the human growth hormone or globin genes can be used as well. Stable cell lines carrying a gene of interest integrated into the chromosomes can also be selected upon co-transfection with a selectable marker such as gpt, G418 or hygromycin. It is advantageous to use more than one selectable marker in the beginning, e.g., G418 plus methotrexate.

The plasmid pC4 is digested with the restriction enzymes and then dephosphorylated using calf intestinal phosphates by procedures known in the art. The vector is then isolated from a 1% agarose gel. The DNA sequence encoding the *B. burgdorferi* polypeptide is amplified using PCR oligonucleotide primers corresponding to the 5' and 3' sequences of the desired portion of the gene. A 5' primer containing a restriction site, a Kozak sequence, an AUG start codon, and nucleotides of the 5' coding region of the *B. burgdorferi* polypeptide is synthesized and used. A 3' primer, containing a restriction site, stop codon, and nucleotides complementary to the 3' coding sequence of the *B. burgdorferi* polypeptides is synthesized and used. The amplified fragment is digested with the restriction endonucleases and then purified again on a 1% agarose gel. The isolated fragment and the dephosphorylated vector are then ligated with T4 DNA ligase. *E. coli* HB101 or XL-1 Blue cells are then transformed and bacteria are identified that contain the fragment inserted into plasmid pC4 using, for instance, restriction enzyme analysis.

Chinese hamster ovary cells lacking an active DHFR gene are used for transfection. Five μ g of the expression plasmid pC4 is cotransfected with 0.5 μ g of the plasmid pSVneo using a lipid-mediated transfection agent such as LipofectinTM or LipofectAMINETM (Life Technologies Gaithersburg, MD). The plasmid pSV2-neo contains a dominant selectable marker, the *neo* gene from *Tn5* encoding an enzyme that confers resistance to a group of antibiotics including G418. The cells are seeded in alpha minus MEM supplemented with 1 mg/ml G418. After 2 days, the cells are trypsinized and seeded in hybridoma cloning plates (Greiner, Germany) in alpha minus

MEM supplemented with 10, 25, or 50 ng/ml of methotrexate plus 1 mg/ml G418. After about 10-14 days single clones are trypsinized and then seeded in 6-well petri dishes or 10 ml flasks using different concentrations of methotrexate (50 nM, 100 nM, 200 nM, 400 nM, 800 nM). Clones growing at the highest concentrations of methotrexate are then transferred to new 6-well plates containing even higher concentrations of methotrexate (1 μ M, 2 μ M, 5 μ M, 10 mM, 20 mM). The same procedure is repeated until clones are obtained which grow at a concentration of 100-200 μ M. Expression of the desired gene product is analyzed, for instance, by SDS-PAGE and Western blot or by reversed phase HPLC analysis.

6. Immunization and Detection of Immune Responses

6(a). *B. burgdorferi* propagation

B. burgdorferi sensu stricto isolate B31 is propagated in tightly-closed containers at 34°C in modified Barbour-Stoermer-Kelly (BSKII) medium (Barbour, A.G., *Yale J. Biol. Med.* 57:521-525 (1984)) overlaid with a 5%O₂/5%CO₂/90%N₂ gas mixture. Cell densities of these cultures are determined by darkfield microscopy at 400X.

Immunization of Mice and Challenge with B. burgdorferi. For active immunizations BALB/cByJ mice (BALB, Jackson Laboratories) are injected intraperitoneally (i.p.) at week 0 with 20 g of recombinant borrelial protein, or phosphate-buffered saline (PBS), emulsified with complete Freund's adjuvant (CFA), given a similar booster immunization in incomplete Freund's adjuvant (IFA) at week 4, and challenged at week 6. For challenge *B. burgdorferi* are diluted in BSKII from exponentially-growing cultures and mice are injected subcutaneously (s.c.) at the base of the tail with 0.1 ml of these dilutions (typically 10³-10⁴ borreliae; approximately 10-100 times the median infectious dose). Borreliae used for challenge are passaged fewer than six times *in vitro*. To assess infection, mice are sacrificed at 14-17 days post-challenge, and specimens derived from ear, bladder, and tibiotarsal joints are placed in BSKII plus 1.4% gelatin, 13 g/ml amphotericin B, 1.5 g/ml phosphomycin, and 15 g/ml rifampicin, and borrelia outgrowth at two or three weeks is quantified by darkfield microscopy. Batches of BSKII are qualified for infection testing by confirming that they supported the growth of 1-5 cells of isolate B31. In some instances seroconversion for protein P39 reactivity is also used to confirm infections (see below). Others have previously shown that mice elicited antibodies to P39 when inoculated with live borreliae by syringe or tick bite, but not with killed borreliae (Simpson, W.J., *et al.*, *J. Clin. Microbiol.* 29:236-243 (1991)).

6(b). Immunoassays

Several immunoassay formats are used to quantify levels of borrelia-specific antibodies (ELISA and immunoblot), and to evaluate the functional properties of these antibodies (growth inhibition assay). The ELISA and immunoblot assays are also used to detect and quantify antibodies elicited in response to borrelial infection that react with specific borrelial antigens. Where antibodies to certain borrelial antigens are elicited by infection this is taken as evidence that

the borrelial proteins in question are expressed *in vivo*. Absence of infection-derived antibodies (seroconversion) following borrelial challenge is evidence that infection is prevented or suppressed. The immunoblot assay is also used to ascertain whether antibodies raised against recombinant borrelial antigens recognize a protein of similar size in extracts of whole borreliae. Where the natural protein is of similar, or identical, size in the immunoblot assay to the recombinant version of the same protein, this is taken as evidence that the recombinant protein is the product of a full-length clone of the respective gene.

Enzyme-Linked Immunosorbant Assay (ELISA). The ELISA is used to quantify levels of antibodies reactive with borrelial antigens elicited in response to immunization with these borrelial antigens. Wells of 96 well microtiter plates (Immunlon 4, Dynatech, Chantilly, Virginia, or equivalent) are coated with antigen by incubating 50 l of 1 g/ml protein antigen solution in a suitable buffer, typically 0.1 M sodium carbonate buffer at pH 9.6. After decanting unbound antigen, additional binding sites are blocked by incubating 100 l of 3% nonfat milk in wash buffer (PBS, 0.2% Tween 20, pH 7.4). After washing, duplicate serial two-fold dilutions of sera in PBS, Tween 20, 1% fetal bovine serum, are incubated for 1 hr, removed, wells are washed three times, and incubated with horseradish peroxidase-conjugated goat anti-mouse IgG. After three washes, bound antibodies are detected with H_2O_2 and 2,2'-azino-di-(3-ethylbenzthiazoline sulfonate) (Schwan, T.G., *et al.*, *Proc. Natl. Acad. Sci. USA* 92:2909-2913 (1985)) (ABTS®, Kirkegaard & Perry Labs., Gaithersburg, MD) and A_{405} is quantified with a Molecular Devices, Corp. (Menlo Park, California) Vmax™ plate reader. IgG levels twice the background level in serum from naive mice are assigned the minimum titer of 1:100.

6(c). *In Vitro* Growth Inhibition Assay

Unlike other bacteria, borreliae can be killed by the binding of specific antibodies to their surface antigens. The mechanism for this *in vitro* killing or growth-inhibitory effect is not known, but can occur in the absence of serum complement, or other immune effector functions. Antibodies elicited in animals receiving immunizations with specific borrelial antigens that result in protection from borrelial challenge usually will directly kill borreliae *in vitro*. Thus, the *in vitro* growth inhibition assay also has a high predictive value for the protective potency of the borrelial antibodies, although exceptions, such as antibodies against OspC which are weak at *in vitro* growth inhibition, have been observed. Also, this assay can be used to evaluate the serologic conservation of epitope binding protective antibodies. A microwell antibody titration assay (Sadziene, A., *et al.*, *J. Infect. Dis.* 167:165-172 (1993)) is used to evaluate the growth inhibition (GI) properties of antisera against recombinant borrelial antigens against the homologous B31 isolate, and against various strains of borrelia. Briefly, 10^5 borrelia in 100 l BSKII are added to serial two-fold dilutions of sera in 100 l BSKII in 96-well plates, and the plates are covered and incubated at 34°C in a 5%O₂/5%CO₂/90%N₂ gas mixture for 72 h prior to quantification of borrelia growth by darkfield microscopy.

6(d). Sodiumdodecylsulfate-Polyacrylamide Gel Electrophoresis (SDS-PAGE) and Immunoblotting

Using a single well format, total borrelial protein extracts, recombinant borrelial antigen, or recombinant P39 samples (2 g of purified protein, or more for total borrelial extracts) are 5 boiled in SDS/2-ME sample buffer before electrophoresis through 3% acrylamide stacking gels, and resolving gels of higher acrylamide concentration, typically 10-15% acrylamide monomer. Gels are electro-blotted to nitrocellulose membranes and lanes are probed with dilutions of antibody to be tested for reactivity with specific borrelial antigens, followed by the appropriate secondary antibody-enzyme (horseradish peroxidase) conjugate. When it is desirable to confirm 10 that the protein had transferred following electro-blotting, membranes are stained with Ponceau S. Immunoblot signals from bound antibodies are detected on x-ray film as chemiluminescence using ECL™ reagents (Amersham Corp., Arlington Heights, Illinois).

6(e). Detection of *Borrelia* mRNA expression

Northern blot analysis is carried out using methods described by, among others, 15 Sambrook *et al.*, *supra*, to detect the expression of the *B. burgdorferi* nucleotide sequences of the present invention in animal tissues. A cDNA probe containing an entire nucleotide sequence shown in Table 1 is labeled with ^{32}P using the *rediprime*™ DNA labeling system (Amersham Life 20 Science), according to manufacturer's instructions. After labeling, the probe is purified using a CHROMA SPIN-100™ column (Clontech Laboratories, Inc.), according to manufacturer's 25 protocol number PT1200-1. The purified labeled probe is then used to detect the expression of *Borrelia* mRNA in an animal tissue sample.

Animal tissues, such as blood or spinal fluid, are examined with the labeled probe using 30 ExpressHyb™ hybridization solution (Clontech) according to manufacturer's protocol number PT1190-1. Following hybridization and washing, the blots are mounted and exposed to film at 70 C overnight, and films developed according to standard procedures.

The disclosure of all publications (including patents, patent applications, journal articles, laboratory manuals, books, or other documents) cited herein are hereby incorporated by reference 35 in their entireties.

The present invention is not to be limited in scope by the specific embodiments described herein, which are intended as single illustrations of individual aspects of the invention. Functionally equivalent methods and components are within the scope of the invention, in addition to those shown and described herein and will become apparent to those skilled in the art from the foregoing description and accompanying drawings. Such modifications are intended to fall within the scope of the appended claims.

Provisional Application Serial No. 60/057,483 filed 3 September 1997 is incorporated by reference herein in its entirety.

TABLE 1. Nucleotide and Amino Acid Sequences

f101.aa

MSKIFLLFNAGFFFLLKIIYVFSYPEIKNFSRQDPVFSIDLKIKVLKYNKKQHIPLFYFSYKVKKGDTFFKIANKING
WQSGIATINLLDSPAVSVGQEIILIPSKKGVFVFDSDKDYRFNNLLLATRDLAKAEKVKIKRNDRVYEFYFFDFVKNP
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t101.aa

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LIPSKKGVFVFDSDKDYRFNNLLLATRDLAKAEKVKIKRNDRVYEFYFFDFVKNPDFGLFSGTELLFFL NANFIFP
LKKFIVSSDFGRNDPFTGNKSFHTGIDLAAPMNAEVYLLLLE

f101.nt

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t101.nt

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f11.aa

VKKYIKTIFLISMVYFYCCTTIKINHDYETDFKVLESPSKYINIDVIKATNEYIYIQITNNSLDVVKINWQNTSLN
NDKIVLKEDLTINNETGYKNKYREFFIGPKTSFKFKVYPLKIHSKNNSNNLSSTIKYPSIFKLNITKVGIEAKK
TINVLITRTTKINITNK

t11.aa

CCTTIKINHDYETDFKVLESPSKYINIDVIKATNEYIYIQITNNSLDVVKINWQNTSLNNDKIVLKEDLTINNET
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f11.nt

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TABLE 1. Nucleotide and Amino Acid Sequences

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t11.nt

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TGA

f12.aa

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GFSVFNTVYLFGNKSSSEDSSFLDFDFNSVYNSGKKPYIRNGYLTYFFAENLAPSINKDYLFDIYANLGFYSG
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NRKTKK

t12.aa

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f12.nt

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TABLE 1. Nucleotide and Amino Acid Sequences

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t12.nt

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TABLE 1. Nucleotide and Amino Acid Sequences

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f129.aa

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t129.aa

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f129.nt

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TABLE 1. Nucleotide and Amino Acid Sequences

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t129.nt

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f142.aa

MDKISILYTLINIIIMLILISIVYLCKRKNSFTKRVFIALAIGIVFGMTIQYFYGTNSEITNETINWISILGDGY
VRLLKMIPIIPLIITSIISAIKLTNSKDVGKMSLLVILTLVFTAGIAAIIGIFTALALGLTAEGLQAGTIEILQSE
KLQKGLEILNQTTITKKITDLIPQNIFEDFAGLRKNSTIGVVIFSAIIGIAALKTSIKKPESIEFFKKIILTLQDI
ILGVVTLILKLPYAILALMTKITAETSEIKSIIKLGEFVIASYIAIGLTFLMHMTLIAINKLNPITFIKKIFPALS
FAFISRSSATIPINIEIQTKNLGVSEGIANLSSSGTSIGQNGCAALHPAMLAIMIAPTQGINPTDISFILTLLIG
LIIITSFGAAGAGGGATTASLMVLSAMNFPVGLVGLVISVEPIIDMGRNAVNVGGAGVISAKQLQFNHNQIYN
QKELVNK

t142.aa

CKRKNSFTKRVFIALAIGIVFGMTIQYFYGTNSEITNETINWISILGDGYVRLLKMIPIIPLIITSIISAIKLTN
SKDVGKMSLLVILTLVFTAGIAAIIGIFTALALGLTAEGLQAGTIEILQSEKLQKGLEILNQTTITKKITDLIPQN
IFEDFAGLRKNSTIGVVIFSAIIGIAALKTSIKKPESIEFFKKIILTLQDIILGVVTLILKLPYAILALMTKITA
TSEIKSIIKLGEFVIASYIAIGLTFLMHMTLIAINKLNPITFIKKIFPALSFAFISRSSATIPINIEIQTKNLGV
SEGIANLSSSGTSIGQNGCAALHPAMLAIMIAPTQGINPTDISFILTLLIGLIIITSFGAAGAGGGATTASLMVLS
AMNFPVGLVGLVISVEPIIDMGRNAVNVGGAGVISAKQLQFNHNQIYNQKELVNK

f142.nt

TAAGAGGTATAATGGATAAAATAAGTATATTATACATTAATCAATATTATAATAATGCTTATTCTAATAAGCA
TAGTTTATCTTGAAAGAAAAATGTTCTTTACAAAAGAGTGTATAGCGTTAGCAATCGGAATAGTATT
TGGAAATGACCATTCAATATTTATGGAACAAATTGAGAAATAACAAACGAAACTATAAATTGATAAGTATTG
GGCGATGGATACGTAAAGCTCCTTAAATGATTATAATCCCCTAATAATAACATCAATAATCTCTGCAATAATAA
AACTAACCAATAGTAAAGATGTTGGAAAATGAGCCTACTTGTATATAACACTAGTATTACAGCAGGTATTGC
TGCCATAATTGGCATTTCACTGCTTAGCATTGGGATTAACAGCGAAGGACTACAAGCGGAACCATCGAAATT
TTACAAAGTAAAAATTGCAAAAGGGCTTGAAATATTAAATCAAACAACAATCACAAAAAAATCACAGATCTTA
TTCCACAAAATATATTGAAGATTGCAAGGGCTTAGAAAAAACTCAACCATCGGGCTGTATATTTCAGCTAT
CATAGGAATAGCCGCCCTAAACATCTACAAAAAGCCAGAACATCAATAGAATTAAAAATAATATTAA
CTCCAAGACATAATATTAGGTGTAGTAATTGATTAAACTAACGCCATTGCTATATTAGCTTAAATGACAA
AAATTACAGCAACCAGCAGAAATCAAAGCATAATAAAGCTTGGAGAATTGTAATTGCTTCTACATTGCATAGG
TCTTACATTCTTATGCATATGACATTAAATTGCAATAAAATAAAATTAAACCAATTACTTTATAAAAAAAATATT
CCAGCACTATCATTGCATTCACTAGGTGAGTGCTGCAACCACATACCAATTAGAAATTCAAACACTAAAA
ATCTGGAGTAAGCGAAGGAATAGCAAATTATCAAGCTCCTTGGAAACATCAATTGGCAGGAAATGGTTGCGAGC
ACTACACCCCGCTATGCTTGCATAATGATAGCAGCAACTCAGGGAAATAACACCCACAGATATTCAATTACTC
ACACTTATTGGATTAATAATAACTTCATTGGAGCTGCTGGCGCTGGTGGAGGCGCAACACAGCCTCACTAA
TGGTGCCTCAGCAATGAACATTCCAGTGGATTGGTAGGACTTGTAAATTCTGTTGAGCCTATAATTGACATGGG
AAGAACAGCTGTTAATGTAGGCGGCTCAATGCTTGCAGGCGTTATATCTGCTAAACAGCTCAAACAAATTCAACC
AATATATACAACCAAAAGAGCTTGTAAACAAATAA

t142.nt

TABLE 1. Nucleotide and Amino Acid Sequences

TGTAAAAGAAAAATGTTCTTTACAAAAGAGTGTATAGCGTTAGCAATCGGAATAGTATTGGAATGACCA
 TTCATATTATGGAACAAATCAGAAATAACAAACGAAACTATAAATTGGATAAGTATTGGCGATGGATA
 CGTAAGGCTCCTTAAATGATTATAATCCCTTAATAACATCAATAATCTCTGCAATAATAAAACTAACCAAT
 AGTAAAGATGTTGGAAAATGAGCCTACTGTAAATTAAACACTAGTATTACAGCAGGTATTGCTGCCATAATTG
 GCATTTCACTGCTTAGCATTGGATTAAACAGCCGAAGGACTACAAGCGGGACCAGTCAAATTACAAAGTGA
 AAAATTGCAAAAGGCCTGAAATATTAAACAAACAATCACAAAAAAATCACAGATCTATTCCACAAAAT
 ATATTGAAAGATTTGCAGGGCTTAGAAAAAACTCAACCATCGGGCTGTGATATTTCAGCTATCATAGGAATAG
 CGCCCTTAAACATCTATCAAAAAGCCAGAATCAATAGAATTTTAAAAAATAATTAAACACTCCAAGACAT
 AATATTAGGTGTAGTAACTTGATTAAACTAACGCCTATGCTATTAGCTTAATGACAAAAATTACAGCA
 ACCAGCGAAATCAAAGCATAATAAAGCTGGAGAATTGTAATTGCTCCTACATTGCCATAGGTCTTACATTTC
 TTATGCATATGACATTAATTGCAATAATAAATTAAACCCAAATTACTTTATAAAAAAAATATCCCAGCACTATC
 ATTTGCATTCAATCTAGGTGAGTGCTGCAACCATAACCCATTAAATAGAATTCAAACAAAAATCTGGGAGTA
 AGCGAAGGAATAGCAAATTATCAAGCTCCTTGGACATCAATTGGCAAAATGGTGTGCAGCACTACACCCCG
 CTATGCTTGCATAATGATAGCACCAACTCAGGGATAAAACCCACAGATATTCAATTACTCACACTATTGG
 ATTAATAATAACTCATTGGAGCTGCTGGCGCTGGTAGGACTGTAAATATCTGTTGAGCCTATAATTGACATGG
 GCAATGAACTTCCAGTGGATTGGTAGGACTGTAAATCTGCTAAACAGCTCAAACAAATTCAACCATAATATACAA
 TTAATGTAGGCGGCTCAATGCTGCAGGCGTTATATCTGCTAAACAGCTCAAACAAATTCAACCATAATATACAA
 CCAAAAGAGCTTGTAAACAAATAA

f147.aa

MKIIIIIGTSAGTSAAKANRLNKLDITIYEKTNIVSFGTGCLPYFVGFFFDNPNTMISRTQEEFEKTGISVKTN
 HEVIKVDAKNNTIVIKNQKTGTIFNNTYDQLMIATGAKPIIPPINNINLENFHTLKNLEDQKIKKLMREEIKNI
 VIIGGGYIGIEMVEAAKRNKRKNVRILQLDKHLIDSFDEEIVTIMEEELTKKGVNLTNEFVKSLIGEKKAEGVVT
 NKNTYQADAVILATGIKPDTEFLENQLKTTKNGAIIVNEYGETSIKNIFSAGDCATIYNIVSKNEYIPLATTANK
 LGRIVGENLAGNHTAFKGTLGSASIKILSLEAARTGLTEKDAKKLQIKYKTIFVKDNHTNYPQEDLYIKLIYE
 ENTKIILGAQAIKGNGAVIRIHALSIAIYSKLTTELGMMDFSYSPPFSRTWDILNIAGNAAK

t147.aa

AAAKANRLNKLDITIYEKTNIVSFGTGCLPYFVGFFFDNPNTMISRTQEEFEKTGISVKTNHEVIKVDAKNNTIV
 IKNQKTGTIFNNTYDQLMIATGAKPIIPPINNINLENFHTLKNLEDQKIKKLMREEIKNIVIIGGGYIGIEMVE
 AAKRNKRKNVRILQLDKHLIDSFDEEIVTIMEEELTKKGVNLTNEFVKSLIGEKKAEGVVTNKNTYQADAVILAT
 GIKPDTEFLENQLKTTKNGAIIVNEYGETSIKNIFSAGDCATIYNIVSKNEYIPLATTANKLGRIVGENLAGNHT
 AFKGTLGSASIKILSLEAARTGLTEKDAKKLQIKYKTIFVKDNHTNYPQEDLYIKLIYEENTKIILGAQAIKG
 NGAVIRIHALSIAIYSKLTTELGMMDFSYSPPFSRTWDILNIAGNAAK

f147.nt

ATGAAAATAATAATTATTGGGGCACATCAGCAGGAACTAGTCGGCAGCTAACAGCAAAACCGCTAAACAAAAAGC
 TAGACATTACTATCTATGAAAAACAAATATTGTATCTTTGGAACCTGTCGGCTGCCCTACTTTGTGGGGGATT
 CTTTGACAACCCAAATACAATGATCTCAAGAACACAAGAAGATTGAAAAAAACTGGAATCTCTGTTAAACTAAC
 CACGAAGTTATCAAAGTAGATGCAAAACAAATACAATTGTAATAAAATCAAAACAGGAACCATTTTAACA
 ATACTTACGATCAACTTATGATAGCAACTGGTCAAAACCTATTATTCCACCAATCAATAATATCAATCTAGAAAA
 TTTTCATACTCTGAAAAATTAGAACAGCGTCAAAAAAATAAAATTAATGGATAGAGAAGAGATTAAATATA
 GTGATAATTGGTGGGATACATTGAATTGAAATGGTAGAAGCAGCAAAATAAAAGAAAAATGTAAGATTAA
 TTCAACTAGATAAGCACACTCATAGATTCTTGAAGAAATGTCACAATAATGGAAGAAGAAGACTAACAAA
 AAAGGGGTTAATCTCATACAAATGAGTTGAAAAAGTTAATAGGAGAAAAAGGCAGAAGGAGTAGTAACA
 AACAAAAAACTTATCAAGCTGACGCTGTATACTTGCTACCGAATAAAACCTGACACTGAATTTTAGAAAACC
 AGCTTAAACTACTAAAATGGAGCAATAATTGTAATGAGTATGGCGAAACTAGCATAAAAATTTCTGC
 AGGAGATTGTCAACTATTATAATAGTAAAGTAAAAAAATGAATACATACCCCTGGCAACACAGCCAACAAA
 CTTGGAAGAATAGTTGGTAAAAATTAGCTGGGAATCATAACAGCATTAAAGGCACATTGGGCTCAGCTCAATT
 AAATACTATCTTACAGAGCTGCAAGAACAGGACTTACAGAAAAAGATGCAAAAGCTCCAAATAAAATATAAAAC
 GATTTTGTAAAGGACAAAATCATACAAATTATTATCCAGGCCAAGAAGATCTTATATTAAATTATGAG
 GAAAATACCAAAATAATCCTGGGCACAAGCAATAGGAAAAATGGAGCCGAATAAGAATTGCTTATCAA

TABLE 1. Nucleotide and Amino Acid Sequences

TTGCAATCTATTCAAAACTTACAACAAAAGAGCTAGGGATGATGGATTCTCATATTCCCCACCCCTCTCAAGAAC
TTGGGATATATTAAATATTGCTGGCAATGCTGCCAATAG

t147.nt

GCCGCAGCTAAAGCAAACCGCTAAACACAAAAGCTAGACATTACTATCTATGAAAAAACAAATATTGTATCTTTG
GAACCTGTGGCCTGCCTACTTGTGGGGGATTCTTGACAACCCCAATACAATGATCTCAAGAACACAAGAAGA
ATTGCAAAAACAGGAATCTCTGTTAAAACCAACACGAAGTTATCAAAGTAGATGCAAAAAACAATACAATTGTA
ATAAAAATCAAAAACAGGAACCATTAAACAAATACTTACGATCAACTTATGATAGCAACTGGTGCAAAACCTA
TTATTCCACCAATCAATAATATACTAGAAAATTTCATACTCTGAAAATTAGAAGACGTCAAAAAATAAA
AAAATTAAATGGATAGAGAAGAGATTAAAATATAGTATAATTGGTGGGATACATTGAATTGAAATGGTAGAA
GCAGCAAAAATAAAAGAAAAATGTAAGATTAATTCAACTAGATAAGCACATACTCATAGATTCTTGACGAAG
AAATAGTCACAATAATGGAAGAAGAACTAACAAAAAGGGGTTAATCTCATACAAATGACTTGTAAAAAGTT
AATAGGAGAAAAAAAGGCAGAAGGAGTAGTAACAAACAAAATACTTATCAAGCTGACGCTGTATACTGCTACC
GGAATAAAACCTGACACTGAATTAGAAAACCAGCTAAAACACTAAAGCAATAATTGTAATGAGT
ATGGCGAAACTAGCATAAAAATTTTCTGAGGAGATTGTGCAACTATTATAATATAGTAAGTAAAAAAA
TGAATACATACCCTGGCAACACAGCCAAACAAACTTGGAGAAATAGTTGGTAAATAGCTGGGATCATA
GCATTAAAGGCACATTGGCTCAGCTCAATTAAAATACTATCTTAAAGCTGCAAGAACAGGACTACAGAAA
AAGATGCAAAAAGCTCAAATAAAATATAAAAGATTAAAGCTTGTAAAGGACAAAATCATACAAATTATTATCCAGG
CCAAGAAGATCTTATATTAAATTATGAGGAAAATACCAAAATACTTGGGGCACAAGCAATAGGAAAA
AATGGAGCCGTAAAGAATTCAATGCTTATCAATTGCAATCTTACAAACAAAAGAGCTAGGGATGA
TGGATTCTCATATTCCCCACCCCTCTCAAGAACTTGGATATATTAAATATTGCTGGCAATGCTGCCAATAG

f152.aa

MLKFEFSDRFLFSYFVLIMFIGSLLMLPISWEGDGKLAYIDALFTAVSAVSITGLTTVKMEGFSTFGFILIMLL
IQLGLGFISITFYLLIPKKKMNLTDARIIKQYSLSNIEYNPIRILKSILFITFSIEMIGLILILICFKLRGVNI
SFLEALFTTISAFCNAGFSMHSESIYAWRDVPEAIVVVSILIIICGGLGFMVYRDVNNTIKNKKLSSLHAKIVFSL
FFLIIIGAILFFFTEMHKLKAGYSMSTLIFNSIFYSISTRAGFNYLDNSLISGRTQIISLPFMFIGGAPGSTAGG
IKITFFFLIVLAVVKNQNGNGYIIGSYKVSIDSIRFALLFFARAIFILSFSSFFMFFFEGGSGNWKVIDLGYEVFS
AFGTVGLSVGTQDLSFWGKVIIIFTMFAGRIGLFSMAVFVSRKSRFEETRPRQDILVG

t152.aa

WEGDGKLAYIDALFTAVSAVSITGLTTVKMEGFSTFGFILIMLLIQLGLGFISITFYLLIPKKKMNLTDARIIK
QYSLSNIEYNPIRILKSILFITFSIEMIGLILILICFKLRGVNISFLEALFTTISAFCNAGFSMHSESIYAWRDV
EAIVVVSILIIICGGLGFMVYRDVNNTIKNKKLSSLHAKIVFSLFFFTEMHKLKAGYSMSTLIFNS
IFYSISTRAGFNYLDNSLISGRTQIISLPFMFIGGAPGSTAGGIKITFFFLIVLAVVKNQNGNGYIIGSYKVSID
SIRFALLFFARAIFILSFSSFFMFFFEGGSGNWKVIDLGYEVFSAGTVGLSVGTQDLSFWGKVIIIFTMFAGRI
GLFSMAVFVSRKSRFEETRPRQDILVG

f152.nt

ATGTTGAAATTGAAATTAGCAGACAGGTTTTACTTTAGTTATTGTTTAATTATGTTTATAGGCTCTCTT
TGTTGATGTTGCCTATTCTGGAGGTGATGGCAAATTAGCATACATTGATGCTTTTACTGCTGTTCTGC
TGTAAGTATTACGGGCTTACAACGGTTAAAATGGAAGGCTTTCTACTTTGTTATTGATAATGTTGCTA
ATCCAGCTGGGGACTGGATTATAAGTATTACTACTTTTATTGCTTATACCTAAAAGAAAATGAATTAA
CAGATGCAAGAATAATAAGCAGTATTCCCTTCAAAATATAGAATATACTTAAAGCATT
GTTTATAACTTTCAATTGAAATGATAGGTTTAATTAAATACTTATTGTTAAACTTAGGGAGTGAATT
TCATTCTTAGAGGTTGTTACGACAATTCTGCTTTGCAATGCAGGTTTCCATGCATTCTGAGAGTATT
ATGCATGGCGAGATGTTCTGAAGCTATAGTTGTGGCTCTATTAAATAATTGTTGGTGGCTGGTTATGGT
CTATAGAGATGTAATAAACACTATTAAAACAAAAAAACTATCGCTTCATGCCAAGAGTATT
TTCTTTTAATTATAATTGTTGCAATTATTGTTTACAGAGATGCATAAAATTAAAAGCTGGTTATTCAATGA
GCACTTTAATTATAATTCAATTGTTTATTGCAATTAGTACAGAGACAGCTGGTTAATTATCTTGTAAATTCTT
AATAAGCGGAAGAACTCAAATAATTCTCACCATTGTTATTGGTGGTGCACCCGGATCAACTGCAGGAGGG
ATTAAGATTACAACATTGTTAATTGTTATTGGCTGTTAAAATCAAAACGGCAATGGATATTATTGGTT

TABLE 1. Nucleotide and Amino Acid Sequences

CTTACAAGGTTCAATAGATAGTATAAGATTGCACTTTATTTTGCAAGAGCTATTTTATTTAAGTTTTC
 TTTTTCATGCTCTTTGGAGGGAGGATCTGGCAATTGAAAGGTATTGATTAGGTTATGAAGTATTTCT
 GCTTTGGAACGGTGGCTTCAGTTGGAGTAACCTCAGGATTGTCATTGGGGAAAAGTCATTATAATTTA
 CTATGTTGCAGGACGAATAGGGCTTTCAATGGCTGTTCAAGAAAGTCGCGTTGAAGAATTAC
 AAGGCCAAGGCAAGATATTTGGTTGGTGA

t152.nt

TGGGAAGGTGATGGCAAATTAGCATACTGATGCTTTTACTGCTGTTCTGCTGTAAGTATTACGGGCCTTA
 CAACGGTTAAATGGAAGGCCTTCTACTTTGGATTATTTGATAATGTCATACTCCAGCTGGGGACTTGG
 ATTTATAAGTATTACTACTTTATTTGCTTACCTAAAAAGAAAATGAATTAAACAGATGCAAGAATAATAAG
 CAGTATTCCCTTCAAATATAGAATATAATCCTATTAGAATTAAAAGCATATTGTTATAACTTTCAATTG
 AAATGATAGGTTAATATTAATACTTATTGTTAAACTTAGGGAGTGAATATTCAATTCTAGAGGCTTGT
 TAGCACAATTCTGCTTTGCAATGCAGGTTTCCATGCATTCTGAGAGTATTGATGGCGAGATGTTCT
 GAAGCTATAGTGTGGCTCTATTAAATAATTGTTGGTGGGTTATGGTCTATAGAGATGAAATAACA
 CTATTAAAACAAAAAAACTATCGCTTCATGCCAAGATAGTTTTCTTAAGCTTCTTTAATTATAATTG
 TGCAATTATTATTTTACAGAGATGCATAAAATTAAAAGCTGGTTATTCATGAGCATTAAATATTCAATTCA
 ATTATTCTCTACCATCATGTTATTGGTGGTCACCCGGATCAACTGCAGGAGGGATTAAGATTACAACATT
 TTTAATTGATTGGCTGTTAAAATCAAACGCCAATGGATATTATTGTTCTTACAAGGTTCAATAGAT
 AGTATAAGATTGCACTTTATTTGCAAGAGCTATTAAAGTTTCTTCAATGCTCTTTTCAAGGTTCAATAGAT
 TTGAGGGAGGATCTGCAATTGAAAGGTATTGATTAGGTTATGAAGTATTCTGCTTGTGAACGGTTGGTCT
 TTCAGTTGGAGTAACCTCAGGATTGTCATTGGGGAAAAGTCATTATAATTAACTATGTTGCAGGACGAATA
 GGGCTTTTCAATGGCTGTTTGTCAAGAAAGTCGCGTTGAAAGAATTACAAGGCCAAGGCAAGATATT
 TGGTGGTGA

f154.aa

MKINKTFILLFLFTKFSFVQAQANQILTEISPLSILSKNGKGSVYLKVSKSSDYILTLKSSNSDFVFKIYDISNK
 KYITDKVKRRDFKIRLDKNSLYAIYVGTKNENIKFSLTDLDFSISSDSLKAKTSKIEKEDLFFTLKDPVLNL
 AKLKKYVLRIFYKSNIYIAYQLENSDDIKVAEFIEDVGWFNLDSVNRNITNIVNFDSINSKGNLYIAFVTKSGAD
 FASELIVKKFNSRKWIDISPGHIEFNGSLLNISIDLKDRLYLAYLREIRGEYKINLISNMGYGSIWTDVIAHLSK
 GDSNVNNSNIGLISEPFLGIFYNYSKSNNEIKSEFIVNNENAWVNANIPSVYMANFIKGFFDSNFNQIIMSFVSENR
 PIVNICPLKSSRWINISPNEMEGLSADIGLYKNNLFLAFEDNNNVRLIYFKKNWYFLNKENFKSNVKSPQIGI
 YGNQGLVISTLSSNSNELFFTLICQ

t154.aa

NQILTEISPLSILSKNGKGSVYLKVSKSSDYILTLKSSNSDFVFKIYDISNKKYITDKVKRRDFKIRLDKNSLYA
 IYVGTKNENIKFSLTDLDFSISSDSLKAKTSKIEKEDLFFTLKDPVLNLAKLKKYVLRIFYKSNIYIAYQLEN
 SDDIKVAEFIEDVGWFNLDSVNRNITNIVNFDSINSKGNLYIAFVTKSGADFASELIVKKFNSRKWIDISPGHI
 ENFGSLLNISIDLKDRLYLAYLREIRGEYKINLISNMGYGSIWTDVIAHLSKGDSNVNNSNIGLISEPFLGIFYN
 YKSNEIKSEFIVNNENAWVNANIPSVYMANFIKGFFDSNFNQIIMSFVSENRPIVNICPLKSSRWINISPNE
 GLSADIGLYKNNLFLAFEDNNNVRLIYFKKNWYFLNKENFKSNVKSPQIGIYGNQGLVISTLSSNSNELFFTL
 CQ

f154.nt

ATGAAAATAATAAGACATTCTTGTATTTACAAAATTTCTTTGTCAGCTCAAGCAAATCAA
 TATTAACAGAAATTAGTCCTTAAGTATTAAAGCAAAATGGGAAAGGAAGTGTACTTAAAGTTAGCAAATC
 TTCCGATTATATTAAACCTAGATAAGAGTTCAATTCCGATTGTTAAAATTTATGACATTCTTAATAAAA
 AAATATATAACCGATAAAAGTAAAAGAAGAGATTAAAGATTAGATAAAATCTCTTATGCAATAATAT
 ATGTTGGTACTAAAATGAAAACATAAAGTTTCGCTTACAGATTAGATTTCATTTAAGTAGCGATTCCCT
 GAAAGCTAAAACATCTAAGATTGAAAAGAAGATTATTTTACTTTAAAGATTGCCTGTTAAATTAACT

TABLE 1. Nucleotide and Amino Acid Sequences

GCCAAGCTTAAAAATATGTATTAAGGATTATAAAAGCAATATTATATTGCTTATCAGCTAGAAAATAGCGATG
 ATATTAAAGTTGCTGAATTATGAGGATTTGGTTAACCTGATTCTGTTAACAGAAATATTACTAA
 TATAGTTAATTGATTTCATTAAATTCTAAAGGAAATTATATATTGCTTTGTTACGAAATCAGGGCTGAT
 TTTGCCAGCGAGCTTATGTTAAAATTAAATAGTAGAAAATGGATTGATATTAGTCCTGGTCACATAGAAAATT
 TTGGATCTTATTAAATATTAGCATTGATTAAAAGATAGGTTGTTAGCATATTAAAGGAAATTAGGGTGA
 ATATAAAATTAATTAAATCTGAATATGGGTACGGAAGTATTGGACCGATGTAATACATGCTTATTAAAGTAAA
 GGTGATTCTAATGTTAATTCAACATGGTTAATATCTGAACCTTTGGCATTTTATAATTATAAGT
 CAAATAATGAGATTAAATCTGAATTATTGTTAACATGAAAATGCTGGTAAATGCAAATATTCTCTGTTA
 TATGGCAATTAAAGGCTTTGTTAAAGGCTTTGATTCTAATTAAATCAAAATTATGAGTTGTTCTGAAAATAGA
 CCTATTGTAACATTGCTTGTAAAAGTAGTAGATGGATTAAATAAGTCTAATGTTGAAATGGAAGGTTAA
 GTGCTGACATTGGGCTTTATAAAAATAATTGTTTAGCTTTGAGGACAATAATAATGTGAGATTAAATTATT
 TAAGAATAAAATTGGTATTGTTAAAGCTGAGAATTAAAGAGTAATGTTAAAAGCCCTCAGATTGGAATT
 TATGGCAATCAAGGGCTGTAATCTACTTTAAGCTCTAATTCCAATGAATTATTGTTACTTGATTGCAAT
 GA

t154.nt

AATCAAATATTAACAGAAATTAGCCTTAAGTATTAAAGCAAAATGGGAAAGGAAGTGTACTTAAAGTTA
 GCAAATCTCCGATTATATTAAACCCCTAGATAAGAGCTCAAATTCCGATTGTTAAAATTATGACATTTC
 TAATAAAAATATATAACCGATAAAAGTAAAAGAAGAGATTAAAGATTAGATAAAATTCTCTTATGCA
 ATAATATATGTTGGTACTAAAATGAAAACATAAAGTTTCGCTTACAGATTAGATTGTTCAATTAAAGTAGCG
 ATTCCCTGAAAGCTAAAACATCTAAGATTGAAAAGAAGATTATTGTTACTTAAAGATTGCTGTTAAA
 TTTAACTGCCAAGCTTAAAGGCTTAAAGGATTATAAGGAAATATTATATTGCTTACAGCTAGAAAAT
 AGCGATGATATTAAAGTTGCTGAATTATTGAGGATGTTGGTTAACTTGTGATTCTGTTAATAGAAATA
 TTACTAATATAGTTAATTGATTGTTCAATTAACTTAAAGGAAATTATATTGCTTGTGTTACGAAATCAGG
 GGCTGATTGCTGAGCTTAAAGGATTAAAGTAAAGGATTGATATTAGTCCTGGTCACATA
 GAAAATTGATCTTATTAAATATTAGCATTGATTAAAAGATAGGTTGTTAGCATATTAAAGGAAATTAA
 GGGTGAATATAAAATTAAATCTGAATATGGGTACGGAAAGTATTGACCGATGTAATACATGCTTATT
 AAGTAAAGGTGATTCTAATGTTAATTCAACATTGGTTAATATCTGAACCTTTGGCATTTTATAAT
 TATAAGTCAAATAATGAGATTAAATCTGAATTATTGTAACAAATGAAAATGCTGGTAAATGCAAATATTCTT
 CTGTTATATGCCAATTATTAAAGGCTTTTGATTCTAATTAAATCAAATAATTATGAGTTGTTCTGAA
 AAATAGACCTATTGTAACATTGCTTGTAAAAGTAGATGGATTAAATAAGCTCTAATGTTGAAATGGAA
 GGTTAAGTGCTGACATTGGGCTTTATAAAAATAATTGTTAGCTTTGAGGACAATAATAATGTGAGATTAA
 TTTATTAAAGAATAAAATTGGTATTGTTAAAGCTGAGAATTAAAGAGTAATGTTAAAAGCCCTCAGAT
 TGGAATTATGGCAATCAAGGGCTGTAATCTACTTTAAGCTCTAATTCCAATGAATTATTGTTACTTGATT
 TGCCAATGA

f157.aa

MKIFLKIGRGLMVFRLKNDYLALISLLIVSFVGILLIYSSDYNISGSLTKNEYIKQTFWVIIGFFLIFIVG
 KYDLKFVYSMVYPLYFLLILALIFTAFTFGMTVNGARSWIGIWLGGQPSEFGKVVIILTSKFYTEKKGYNEFFTF
 ITAFLLIFPSVILILLQPDFTGTAIVYLTIIFIFISFFAGIDLHYVLAFALIGFFSFVFAILPVWYEYKVNMGNVFYL
 IFSNPFYFRVIMGVLLILLISVLGFFISKYGLSIKIIYFYVFFASSILVSIVFSKVLSKLMKTYQIKRFLVFLD
 PAIDAKGAGWNLNQVKIAIGSGGLLGKFLKGPyTHANYVPSQSTDIFISFISILAEFGFLGVSTILILFFFLFFKFL
 IIMNKSQDRYMLVISGILGLFFHTSFNVGMSLGVLPITGIPFPFLSYGGSSTITFFLAMSFYFNIESIVAMD

t157.aa

RKNYDYLALISLLIVSFVGILLIYSSDYNISGSLTKNEYIKQTFWVIIGFFLIFIVGKYDLKFVYSMVYPLYFLLI
 LALIFTAFTFGMTVNGARSWIGIWLGGQPSEFGKVVIILTSKFYTEKKGYNEFFTFITAFLLIFPSVILILLQPD
 FGTAIVYLTIIFIFISFFAGIDLHYVLAFALIGFFSFVFAILPVWYEYKVNMGNVFYLIFSNPFYFRVIMGVLLIL

TABLE 1. Nucleotide and Amino Acid Sequences

LISVLGFFISKYGLSIKIIYFYVFFASSILLVSIVFSKVLSQLMKTQIKRFLVFLPAIDAKGAGWNLNQVKIAI
GSGGLLGKFLKGPYTHANYVPSQSTDIFISILAEFGFLGVSTILILFFFLLFKFLIMNKSQDRYMALVISGIL
GLLFFHTSFNVGMSLGVLPITGIPFPFLSYGGSSTITFFLAMSPYFNIESIVAMD

f157.nt

ATGAAGATATTCTAAAGGTTAGGCCGTGGTATATTAGGTAGATTAATGGTTTTAGAAAAAAATTATGATTATT
TGGCTTGTATAAGCTTACTTATAGTTCTTGTGGTATATTGGTGTATTATTCTAGCATTATAATATTAGTGG
ATCTTTAACCAAGAATGAATATATAAAAACAAACCTTGTGGTAATTATTGGATTTTCTAATTTTATAGTGGC
AAATATGATTAAACCTTGTGGTATAGCATGGTATACCTTTATATTGGTATTAAATATTGGCTTAATTTTACTG
CATTTTTGGAAAGACAGTAAATGGAGCAAGATCTGGATTGGCATATGGAAACTTGGAGGACAGCCTCTGAATT
TGGTAAAGTTGTATTATTAAACCTTCAAAATTACACTGAAAAAAAGGGTTATAATGAATTTTACCTTT
ATTACTGCATTTTATTAAATTTCATCGGTAACTCTTATATTGCAACCTGATTGGTACAGCAATAGTAT
ATTAAACCATTTTATTTTATTCTTTGCAGGAATAGATTGCACTATGTTAGCATTGCGTTGATAGG
GTTTTTTCTTTGTTTGCATTACCGGTTGGTATGAATATAAGGTGAATATGGTAATGTATTATCTT
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TTCAATAGTGTCTTCAAAGGTTCTTCAAAGTAAATGAAGACTTACAGATTAAACGGTTTTGGTATTCTTAGAT
CCGGCTATTGATGCTAACGGGTGCTGGTGAATTAAACGGTTAAATAGCAATTGGTCTGGCGGTCTTGG
GCAAAGGATTAAAGGGACCTTACCCACGCTAATTATGTGCCATCTCAAAGCACAGATTATTATTCTAT
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ATAATAATGAATAAAAGTCAAGATAGATATGGCCTTAGTAATATCTGGAAATTGGGACTTTATTTTCATA
CTTCTTTAATGTTGGAATGTCTTAGGAGTTCTCCTATTACCGGGATTCCCTTCTCTTCTATGGAGG
TTCTTCTACTATTACATTAGCAATGTCTTTATTAAATATTGAATCAATAGTGTATGGATTGA

t157.nt

AGAAAAAAATTATGATTATTGGCTTGTATAAGCTTACTTATAGTTCTTGTGGTATATTGGTGTATTATTCTA
GCGATTATAATTAGTGGATCTTAAACCAAGAATGAATATATAAAAACAAACCTTGTGGTAATTATTGGATT
TCTAATTAAATTAGTGGCAAATATGATTAAAATTGTGGTATAGCATGGTATACCTTTATATTGGTATTAAATA
TTGGCTTTAATTAACTGCATTGGAATGACAGTAAATGGAGCAAGATCTGGATTGGCATATGGAAACTTG
GAGGACAGCCTCTGAATTGGTAAAGTTGTATTATTAAACCTTCAAAATTACACTGAAAAAAAGGGTTA
TAATGAATTAACTTATTACTGCATTAACTTATTAATTGGTGTGGTAATTCTTATATTGCAACCTGAT
TTGGTACAGCAATAGTATTTAAACCATTTATTTATTCTTTTGCAAGGATAGATTGCACTATGTT
TAGCATTGCGTTGATAGGGTTTTCTTTGCAATTACCGGTTGGTATGAATATAAGGTGAATAT
GGTAATGTATTCTTCTTCAATCCTTTATTTAGAGTAATAATGGAGTGTGCTTAAATTCT
TTGATTCTGTTAGGATTTCATTCTAAATATGGTTGACTATTAAATAATTATTGTATTGG
CAAGTTCTATTAACTAGTGTCTCAAAGGTCTTCAAAGTAAATGAAGACTTACAGATTAAACG
GTTTTGGTATTCTTAGATCCGGCTATTGATGCTAACGGGTGCTGGTTGGAATTAAACGGTTAAATAGCAATT
GGTCTGGCGGTCTTTGGCAAGGATTAAAGGGACCTTACCCACGCTAATTATGTGCCATCTCAAAGCA
CAGATTAAATTCTTCTATTCTGCCGAAGAGTTGGGTTTTGGGTGTTAGCACTATTAAATTATTGG
CCTTTTTAAATTGGATAATAATGAATAAAAGTCAAGATAGATATGGCCTAGTAATATCTGGAAATTGG
GGACTTTATTTCTACTTCTTAAATGTTGGAATGTCTTAGGAGTTCTCCTATTACCGGGATTCCCTTC
CTTTCTCTTATGGAGGTTCTACTATTACATTAGCAATGTCTTTATTAAATATTGAATCAAT
AGTTGCTATGGATTGA

f17.aa

MIVFLFFSIYLIILFKRSSNSPLYFVPDTKFETLSIRIVLSCSLLIFFCTMLDARPSTIAVFPPTPGSPISIALFL
FLLKSIFVRVLISASLPTKGSNFLAFASAVKFLTYFPISKCSFSSRISSNSL

TABLE 1. Nucleotide and Amino Acid Sequences

t17.aa

PLYFVPDTKFETLSIRIVLSCSLLLIFFCMILDARPSTIAVFPTPGSPISIALFLFLLKSIFVRLISASLPTKGS
NFLAFASAVKFLTYFPISKCSFSSRISSNSL

f17.nt

ATGATTGTGTTTGTCCCCAATATACTTAATTATATTAAACGATCTTCAAACCTGCCCTCTATATTTG
TTCCCGATACCAAGTTGAAACCTTAAGCATTAGAATTGTTGTCTTGTAGTTGCTACTTATTTTTTGAC
TATGCTTGATGCAAGGCCTTCAACTATTGCTGTTTCCCACACCAGGTCGCCTATTAGCATTGCACTATTTTA
TTTCTCTCAAGAGTATATTGTAAGAGTTAACCTCTGCTCTTCCAACCAAGGGTCTAATTTGGCTT
TTGCAAGTGTGTTAAATTTGACATACTTCCAATTCAAAGTGCTCATTTCAAGTCGTATTCTTCATCAAA
TTCTTGCTAG

t17.nt

CCTCTATATTTGTTCCCGATACCAAGTTGAAACCTTAAGCATTAGAATTGTTGTCTTGTAGTTGCTACTTA
TTTTTTTTGCACTATGCTTGATGCAAGGCCTTCAACTATTGCTGTTTCCCACACCAGGTCGCCTATTAGCAT
TGCACTATTTTATTCTCTCAAGAGTATATTGTAAGAGTTAACCTCTGCTCTTCCAACCAAGGGGTCT
AATTTTTGGCTTTGCAAGTGTGTTAAATTTGACATACTTCCAATTCAAAGTGCTCATTTCAAGTCGTA
TTTCTTCATCAAATTCTTGCTAG

f170.aa

MKAFKVKNLRRFSNFIRILVIVLFLNSLLSLFVFLAGSYNIFVYNFQKFYLDLAIILSSVSFGLESTRLIFFYFLK
NKKIKYYLILIFSIIFFIALVFKIFLSGNK

t170.aa

YNIFVYNFQKFYLDLAIILSSVSFGLESTRLIFFYFLKNKKIKYYLILIFSIIFFIALVFKIFLSGNK

f170.nt

ATGAAAGCTTTAAAGTAAAAATCTAACGACGTTTCAAATTATTAGAATTGGTTATTGTATTGTTTAA
ATTCTTGTTAAGTTGTTCTGTTGGCTGGTTCTTACAATATTGTTACAATTTCAGAAATTATCT
TGATCTTGCTATTATTTAAGCTCTGTTCTGGACTTGAATCTACTAGACTGATATTGTTATTGAA
AATAAAAAATTAAGTATTATTTAATTAGTTATAATTGCTCTGTTAAAATT
TTCTTGCTGGTAATAA
ATAG

t170.nt

TABLE I. Nucleotide and Amino Acid Sequences

TACAAATTTTTGTTTACAAATTTCAGAATTTATCTTGATCTTGCTATTATTTAAGCTGTGTTCTTTGGAC
 TTGAAATCTACTAGACTGATATTTTATTTGAAAAATAAAAAATAAGTATTATTTAATTTAATTTTAG
 TTTTATAATTTTCTATTCGCTCTGTTTAAATTTCTGGTAATAATAG

f186.aa

MKKLICIFTLFLSQACNLSPMKIDTKEDMKILYSEIAELRKKNLNHLEIDDTLEKVAKEYAIKLGENDRITHTL
 FGTPPMQRHKYDQSFNLTPETLASGIELNPVVAWLNSPSHKEALINTDTDKIGGYRLKTTDNIDIFVVLFGKRK
 YKN

f186.aa

TMKIDTKEDMKILYSEIAELPKKLNLNHLEIDDTLEKVAKEYAIKLGENDRITHTLFGTPPMQRHKYDQSFNL
 TELASGIELNPVVAWLNSPSHKEALINTDTDKIGGYRLKTTDNIDIFVVLFGKRKYKN

f186.nt

ATGAATAAATTGATTATAAATTTACACTGTTTATCTCAAGCATGCAATTAAAGTACAATGCATAAAATAGATA
 CAAAAGAAGATATGAAATACTATATTCAAGAATTGCTGAATTGAGAAAAAAATTAATCTAAACCATCTAGAAAT
 AGATGATACCCCTTGAAATAAGTTCGAAAGAATATGCCATTAAACTGGGAGAAAATAGAACAAATAACTCACACCCCTT
 TTGGCACACCCATTGCAAGATACATAATACGATCAATCCTTAATTAAACAAGAGAAACTGGCATCAG
 GAAATTGAACTTACAGAGTACTTATGCAAGCTTAATAGTCCAAGCCACAAAGAACGCTTATTAAATACAGATA
 CGATAAAATAGCTGGCTATAGATTAAAAACGACTGACAATATAGATATTTGTAGTTCTTTGGAAAAAGAAAA
 TATAAGAATTGA

f186.nt

ACAATGCATAAAATGATACAAAGAAGATATGAAAATTCTATATTCAAGAAATTGCTGAATTGAGAAAAAAATTA
 ATCTAAACCATCTAGAAATAAGATGATACCCCTTGAAAAAGTTGCAAAGAATATGCCATTAAACTGGGAGAAAATAG
 AACAAATAACTCACACCCCTTTGGCACACCCCAATGCAAAGAACATACATAATACGATCAATCCTTAATTAAACA
 AGAGAACATCTGGCATCAGGAATTGAACTTAAACAGAGTAGTTATGCAAGCTTAATAGTCCAAGCCACAAAGAAC
 CTCTTATTAAATACAGATAACCGATAAAATAGCTGGCTATAGATAAAAACGACTGACAATATAGATATTTGTAGT
 TCTTTTGAAAGAAATTAAAGAATTGA

f196.aa

MKLKAPMILLVLLIAAFFISCLFFAGMLINSKLVDQQFNLMINLIESIKSSFNLVISSMEEKVRVSSMYFNSAEK
 PNEASKIKSKPLSFISDQSEILQITGSNMIVTDKEGKIVFTAVKDNDFGKSIGDREYFTKLKESNSIVYNSFVM
 LADPGSIEESILKTIISKIKKKQIPYIILIGPLRDFETDNIIFGYFMFLYSMDIYIYRSFRGINFGILSSGRALAYD
 TTGRLLVHVVLPGDILTDISASYSNIIKTSEDLLQKNKEISTVYYYDPKSNKYYVGISQKVLLNLSNMKFILLM
 RTSEDDFTYMSRATTIILASFVFTLLMLIAATLYLVKKLSSSLNKILEYSERLASGNFTADINFCKWDTVELYSL
 YEGLEQLRITNFSSVANGVIEENLDYIYENAIQIANASQNLSSGAVEQASTLEQMTANIEQISQGVSENTENAATTEK
 IAVNTNEPTKEGHZSVVKAEZAMT/ITEKIGIIDEITRQTNLLALNASIEAARVGEKGKGFEVVAAEVRLKLADQSK
 ESAREIIDIANP.SLTVASPAGENFEQIVPGMEOQTARLVKNISNESYKQSVQIEQFKNAIEQVSQLVQTTASSSEEL
 SAMSERKMLESVYKELKESVVDYFKIEK

TABLE 1. Nucleotide and Amino Acid Sequences

t196.aa

MLINSKLVDQQFNLMINLIESIKSSFNLYISSMEEKVRVSSMYFNSAEKFNEASKIKSKRLSFISDQSEILIQTGS
 NMMVTDKEGKIVFTTAVKDNDFGKSIGDREYFTKLKESNSIVYNSFVMLADPGSIEESLLKDISKIKNKKGQIPY
 ILIGMPLRDFETDNIFYFMFLYSMDIYIYRSFRGINFGILSSGRALAYDTGRLLVHHVLPGDILTDISASYSNI
 IKKTSEDLLQKNKEISTVYYDPKSNKKYVGISQKVLLNLSSNNKFILLMRTSEDDFYYMSRATTIILAISFVFTLL
 MLAIAATLYLVKKLSSSLNKILEYSERLASGNFTADINFOKWDTVELYSLYEQLRTNFSSVAKVIEFLDLYE
 NAIQIANASQNLSSGAVEQASTLEQMTANIEQISQGVSENTENAATTEKIAVNTNERTKEGHKSVVKAIEAMTVIT
 EKIGIIDEITRQTNLALNASIEAARVGEKGKGFEVVAEVRKLADQSKESEI IDIANRSLTVASRAGENFEQI
 VPGMEQTARLVKNISNESYKQSVQIEQFKNAIEQVSQLVQTTASSSEELSAMSEKMLSVKDLKESVDYFIEK

f196.nt

ATGAAGCTTAAAGCTAGGATGTTGCTACTGTTATTCTGATAGCATTCTTATATCAATTGTTTTGCTT
 TTGGAATGCTTATTAATAGTAAATTGGTGGATCAACAGTTAATCTTATGATAAATCTTATTGAAAGCATTAAAG
 TTCTTTAATCTTACATCTCTCAATGGAAGAGAAAGTTAGGGTTAGTCCATGTATTCAACTCTGCTGAAAAAA
 TTTAATGAGGCTAGTAAATTAAACAGGTTGAGCTTATTTCAGATCAATCTGAAATTCTTATTCAAACCG
 GTAGTAATATGATGGTTACAGACAAAGAAGGTAAGTGGTTACTACGGCGGTTAAGGATAATAGTATTGTTGG
 CAAATCTATTGGGGATAGAGAATATTTACAAAAGCTAAGGAGTCAATAGTATTGTTACAATTCTTGTGATG
 TTGGCAGATCCGGGCTATTGAGGAGTCTTACTTAAAGATAATTCAAGATAAAAAAATAAAAAGGTAGATT
 CTTACATATTAAAGGTATGCCATTAAAGAGATTGGTAAACAGATAACATTGTTGTTATTGTTATTCTTATT
 AATGGATTATATATAGGTCTTATAGGGATTAAATTGGAATACTCTAGCGGCGTAGCTTATGAT
 ACTACGGGTAGATTGTTGGTTCATCATGTTAGTATTGCCAGGTGATATTGACTGATATTAGTCTTATTCCA
 ATATTATTAAAGAAAACATCTGAAGATTGTTGCAAAGAATAAAAGAAATTCAACTGTTATTATTATGATCCTAA
 AAGCAATAAGAAATATGTTGGAATTAGTCAAAGGTTATTAAACTTGTCTAATAAAATTATTCTTTAATG
 AGAACTTCAGAGGACGATTTTATTACATGTCACGAGCTACAACATAATCTAGCAATTAGTATTGTTATTAC
 TACTTATGCTTGTCTTGCATTGCAACTCTTATCTGTGAAAAAGTTAAGCTCTTGTGAAATAAGATACTGGAAATT
 TGAGAGACTTGTCTGGTAATTCTACTGCTGATATTAAATTGCAAAATGGGAACTGTAGAGCTTACAGTTG
 TACGAAGGGCTTGAGCAGTTGAGAACCAATTTCAGTTGCAAAAGGAGTTATTGAAATCTAGATTATCTT
 ATGAAAATGCAATTCAAATAGCAAATGCAAGCCAGAATTAAAGTTCTGGCGCTGAGCAGGCTCTACTTT
 GCAAATGACAGCAAATTGAGCAAATTCAAGGTGTTCTGAGAAACTGAAATGCAAGCTACTACTGAAAAAA
 ATTGCTGTTAATACTAATGAAAGGACTAAAGAGGGGCAATACTGTGTTAAGGCTATTGAGGAATGACTGTA
 TTACTGAAAAATTGGAATTATTGATGAGATAACAAGCCAACCAATTGCTTGTCTTAAATGCCCTGATTGAAGC
 TGCACGAGTGGGAGAAAAGGGCAAGGGATTGAAAGTGGTAGCTGCTGAGGTTAGAAAGCTTGAGATCAAAGCAA
 GAATCAGCAAGAGAGATTATTGATATTGCAAACAGAAAGTTAACTGTTGCAAGTCGTGCTGGGAAAATTGAAAC
 AAATAGCTCCTGGTATGGAACAAACAGCCAGACTTGTAAAAAAATTCTAATGAAAGTTATAAGCAAAGTGTCA
 AATAGAGCAATTAAAAATGCAATAGAGCAGGTTAGTCAGTTAGTCAAACACTACAGCCTCAAGCAGTGAAGAGCTT
 TCTGCAATGCTGAAAAGATGTTAGAGAGTGTAAAAGATTAAAGATCTGTTGATTATTAAAGATCGAAAAGT
 AA

t196.nt

ATGCTTATTAATAGTAAATTGGTGGATCAACAGTTAATCTTATGATAAATCTTATTGAAAGCATTAAAGTTCTT
 TTAATCTTACATCTCTCAATGGAAGAGAAAGTTAGGGTTAGTCCATGTATTCAACTCTGCTGAAAAATTAA
 TGAGGCTAGTAAATTAAACAGGTTGAGCTTATTTCAGATCAATCTGAAATTCTTATTCAAACCGGTAGT
 AATATGATGGTTACAGACAAAGAAGGTAAGTGGTTACTACGGCGGTTAAGGATAATAGTATTGTTGGCAAAT
 CTATTGGGGATAGAGAATATTTCAGGACTTAAGGAGTCAATAGTATTGTTACAATTCTTGTGATTGTTGG
 AGATCCGGGCTATTGAGGAGTCTTACTTAAAGATAATTCCAAGATAAAAAAATAAAAAGGTAGATTCTTAC
 ATATTAAATAGGTATGCCATTAAAGAGATTGAAACAGATAACATTGTTGTTATTGTTCTTATTCAATGG
 ATTATATATATAGGTCTTTAGAGGATTAAATTGGAATACTCTAGCGGCGTAGCTTATGATACTAC
 GGGTAGATTGTTGGTTCATCATGTTAGTATTGCCAGGTGATATTGACTGATATTAGTCTTATTCAAATATT

TABLE 1. Nucleotide and Amino Acid Sequences

ATTAAGAAAACATCTGAAGATTGTTGCAAAAGAATAAAGAAATTCAACTGTTATTATTATGATCCTAAAAGCA
 ATAAGAAATATGTGGATTAGTCAAAAGGTGTTAAACTGTCTAATAATAAAATTATTCTTTAATGAGAAC
 TTCAGAGGACGATTATTACATGTCAAGCTACAACATAATCTTAGCAATTAGTTGTATTACATTACTT
 ATGCTTGCTATTGCAACTCTTATCTGTGAAAAGTTAAGCTCTTCTGAAATAAGACTGGAATTCTGAGA
 GACTTGCTCTGGTAATTACTGCTGATATTAAATTGGCAAATGGGAACTGTAGAGCTTACAGTTGTACGA
 AGGGCTTGAGCAGTTGAGAACCAATTTCAGTTGCAAAAGGAGTTATTGAAAATCTAGATTATCTTATGAA
 AATGCAATTCAAATAGCAAATGCAAGCCAGAATTAAAGTTCTGGCGCTGTTGAGCAGGCTTACTTAGAGCAA
 TGACAGCAAATATTGAGCAAATTTCACAAGGTGTTCTGAGAAATACTGAAAATGCAGCTACTGAAAAATTGC
 TGTTAATACTAATGAAAGGACTAAAGAGGGCATAAAATCTGTTGTTAAGGCTATTGAGGCAATGACTGTAATTACT
 GAAAAAATTGAAATTATTGATGAGATAACAAGGCAAACCAATTGCTTAAATGCCCTGATTGAAGCTGCAC
 GAGTGGGAGAAAAGGGCAAGGGATTGAGTGGTAGCTGCTGAGGTTAGAAAGCTTGAGATCAAAGCAAAGAAC
 AGCAAGAGAGATTATTGATATTGAAACAGAACAGTTAAGCTGCAAGTCGTGCTGGGAAAATTGAAACAAATA
 GTTCTGGTATGGAACAAACAGCCAGACTTGTAAAAAAATTCTAATGAAAGTTATAAGCAAAGTGTCAAATAG
 AGCAATTAAAATGCAATAGAGCAGGTTAGTCAGTTAGTCAAACACAGCCTCAAGCAGTGAAGAGCTTCTGC
 AATGCTGAAAAGATGTTAGAGAGTGTAAAAGATTAAAGAATCTGTTGATTATTAAAGATCGAAAAGTAA

f899.aa

MRFIIAFLMILNQGFSNLSPEDIIFESSYEVAIKKAQKLKNVLILVGRDIKENLIKDFLNSFTNGEIIHKVS
 RKSFLVIDKDNEIFNKINLQKSPTIFFVDSKNEQIKAAYVGAVLSSVQFDKDFLNYVMGAIKSTSVLKKQKDYEI
 NTADERTFFYKTLKGDWRLKFNGKDRKLVLFDLKEFLVFKDINENKLYAIPKSIGNIYFSLLGNEEWKLFKGK
 K

t899.aa

f899.nt

ATGAGATTATAATTGCAATTAAATGATTAAATCAAGGATTTCAAATTGTTCTTGCCTCCGAAAGATA
 TTATTGAGAGTTCTTATGAGGTTGCAATTAAAAAGCTAAAAATTGAAATAAAATGTTAATTGGTTGG
 TAGAGATATTAAAGAAAATTAAATAAAAGATTTTAAACTCTTACAAATGGTGAATTATTACAAAGTATCT
 AGAAAAAAGTGTGTTAGTTATTGATAAGGATAATGAAATTAAATAAAATTAAATCTACAAAAAGTCCGACTA
 TTTTTTTGTTGATTCTAAGAATGAGCAAATAAGCAGCTTATGTGGAGCTGTTGAGCAGTGTCAATTG
 TAAGGATTTTAAACTATGTTATGGGAGCTATAAAATCAACAAGTGTAAAGCAAAAGATTATGAAATT
 AATACTGCTGATGAGAGAACCTTTTACAAACATTAAAGGTGATTGGCATTAAAGTTAATGTTAAAGACA
 GAAAGCTTGTCTTTGATAACAGATCTAAAGAATTGTTAGTTAAAGATATTAAATGAAACAAGCTTATGC
 TATTCTAAGTCTAGGATTGTAATTATTCTATTGGAAATGAAGAATGGAAGCTTGTGAAAGATA
 AAATAAA

t899.nt

TTGCCTCCGAAAGATATTATTGAGAGTTCTTATGAGGTTGCAATTAAAAAGCTAAAAATTGAATAAAATG
 TTTAATTGGTTGGTAGAGATATTAAAGAAAATTAAATAAAAGATTTTAAACTCTTACAAATGGTGAAT
 TATTACAAAGTATCTAGAAAAGTGTGTTAGTTATTGATAAGGATAATGAAATTAAATAAAATTAAATCTA
 CAAAAAAGCCGACTATTGTTGATTCTAAGAATGAGCAAATAAGCAGCTTATGTGGGAGCTGTTG
 GCAGTGTCAATTGATAAGGATTAAACTATGTTATGGGAGCTATAAAATCAACAAGTGTAAAGCA
 AAAAGATTATGAAATTAAACTGCTGATGAGAGAACCTTTTACAAACATTAAAGGTGATTGGCATTAAAG
 TTTAATGGTAAAGACAGAACAGCTGTTCTTTGATACAGATCTAAAGAATTGTTAGTTAAAGATATTAAATG
 AAAACAAGCTTATGCTATTCTAAGTCTAGGATTGTAATTATTCTATTGGAAATGAAGAATGGAA
 GCTTTTGAAAATAAAATAA

TABLE 1. Nucleotide and Amino Acid Sequences

f924.aa

MQDRKFSFRKYFLISVFLIFIVSGITYFYSTQMLEKSQKCVEDNLDAKVKLVDMEDFYFDLNECLNMDDFFIPRPD
FLNENLNKNLVDGLIKNKFLDENFFKDLWIKKENLFNVDIEKENEKLIDKILEISK

t924.aa

TQMLEKSQKCVEDNLDAKVKLVDMEDFYFDLNECLNMDDFFIPRPDFLNENLNKNLVDGLIKNKFLDENFFKDLW
IKKENLFNVDIEKENEKLIDKILEISK

f924.nt

ATGCAAGATAGAAAGTTAGTTAGAAAATATTTTAATTTCAGTATTTGATTTTATTGTTCTGGTATTA
CTTAATTCTATTCAACACAAATGTTGAAAAATCTAAAAGTGTGTTGAAGACAATTAGACGCTAAGGTTAAATT
AGTTGATATGGAAGATTTTATTGATTAAATGAATGTCTAAATATGGATGATTTTTATTCCAAGACCTGAT
TTTTAAATGAAAATTAAAGAATTAGTTAGTTGATGGATTGATGAAAATAATTCTTGATGAGAATT
TCAAGGATCTTGGATTAAGGAAAATTATTAAACGTTGATATTGAGAAGGAGAATGAAAATTAAATAGATAA
GATTTAGAAATTCCAATGA

t924.nt

ACACAAATGTTGAAAAATCTAAAGTGTGTTGAAGACAATTAGACGCTAAGGTTAAATTAGTTGATATGGAAG
ATTTTTATTTGATTAAATGAATGTCTAAATATGGATGATTTTTATTCCAAGACCTGATTAAATGAAA
TTTAAATAAGAATTAGTTAGTTGATGGATTGATGAAAATAATTCTTGATGAGAATT
ATTAAAAGGAAAATTATTAAACGTTGATATTGAGAAGGAGAATGAAAATTAAATAGATAAGATTAGAAATT
CCAAATGA

f925.aa

MIRKYLIYISLLFIVFEVYSKPAFISQDDSYELDFSSGEVDISVNTNSKFNLFSKDESWIYIKSIENEAFIKLIGE
SYDNGAVFTFQTFKKKEGKIKLVFTYQNVKDSSEFNKIIILKITKNFEVAIPQGVGGGSSRDNNIETGNNL
ISGATSKIEIIVRALNLSYINDYKGAIIDLNNKYNFNDKYILLKAEIHYKNGDYLKSYENYLKLKSKYFQSIVFDL
RLAIELNIKEEVLENARYLVEKNDSESIYLEIIFEFLVTRGEHEFALNFSSLYFPKYINSSFS
DKYSYLLGKLYESES
SESKHKDFLKALHYYKLVIDNYPFSYYYERAKIRYFLKRFF

t925.aa

KPAFISQDDSYELDFSSGEVDISVNTNSKFNLFSKDESWIYIKSIENEAFIKLIGE
SYDNGAVFTFQTFKKKEGKIKLVFTYQNVKDSSEFNKIIILKITKNFEVAIPQGVGGGSSRDNNIETGNNL
ISGATSKIEIIVRALNLSYINDYKGAIIDLNNKYNFNDKYILLKAEIHYKNGDYLKSYENYLKLKSKYFQSIVFDL
RLAIELNIKEEVLENARYLVEKNDSESIYLEIIFEFLVTRGEHEFALNFSSLYFPKYINSSFS
DKYSYLLGKLYESES
SESKHKDFLKALHYYKLVIDNYPFSYYYERAKIRYFLKRFF

f925.nt

ATGATTAGAAAATATTGATTATATAAGTTGCTATTGTTTGAAGTTACTCTAACGCCAGCTTTATAA
GTCAAGACGATTGAGCTGAGCTGTTAGTAGTGGAGAGGTAGATATTAGTGTAAATACCAATTCAA
TCTTTCTTTAAAGATGAGTCTGGATTATCAAAAGCATTGAAAATGAAGCTTTATTAAGTTAATTGGAGAA

TABLE 1. Nucleotide and Amino Acid Sequences

TCTTATGATAACGGTGCTTTACTTTCAAGACTTTAAAAAGAAGGCAAAATTAAATTGGTTTCACTTATC
 AAAATGTTAAAGATTCAAGTGAATTAAATAAAAATTATCTTGAAGGAACTTCAAAAGAATTGAAAGTTGCAATTCC
 ACAAGGCCTGGTGGCTCTAGCAGGGACAATAACATTGAAACTGGTAATACTTGAACCTGGGGGGGGAGT
 ATTAGCAGGGCAACTCTAAAGAGATTATTGTTAGGGCTTAAATTGCTACATAATGATTACAAAGGAGCAA
 TAGATTGCTTAATAAGTATAATTCAATGACGATAAATATTTATTGAAAGCGGAAATTCAATTATAAAAATGG
 TGATTATTAAATCTTATGAAAATTATTGAAATTGAGAGTAAATATTCAAAAGCATTGTTTGATCTAATT
 AGGCTTGCTATAGAATTAAATTAAGAAGAGGTTAGAGAACGCTAGATATTGAAAGAATGTTGATT
 TTTCTGAGAGCATTATCTGAGATCTTGAATTCTAGTAACAAGGGAGAGCATGAGTTGCTTAAATTGAG
 CTCTCTTACTTCCTAAGTATAATTCAAGCTTCAGACAAATATACTTATTGTTGGAAAACTTATGAG
 TCTGAGAGCAAGCATAAAGATTAAAGGCTTGCAATTACTATAATTGTTATTGATAATTACCCCTTGTAGTT
 ATTATTATGAGAGAGCCAAGATAAGATATTATTAAAGCGTTTTAGT

t925.nt

AAGCCAGCTTTATAAGTCAGACGATTGCTATGAGCTTGAATTAGTAGTGGAGAGGTAGATATTAGTGTAAATA
 CCAATTCAAAATTAAATCTTCTTTAAAGATGAGTCTGGATTATCAGAAAGCATTGAAAGCATTGTTTAT
 TAAGTTAATTGGAGAATCTTATGATAACGGTGCTTTACTTTCAAGCTTAAAGAAGGCAAATTAAA
 TTGGTTTCACTTATCAGAAAGATTCAAGTGAATTAAATAAAATAATTATCTTGAAGGAAATTACAAAGAATT
 TTGAAGTTGCAATTCCACAAGGCGTGGTGGCTCTAGCAGGGACAATAACATTGAAACTGGTAATAATCTTGA
 ACTTGGGGGGGGAGTATTAGCGGGCAACATTCTAAAGAGATTATTGTTAGGGCTTAAATTGCTACATAAT
 GATTACAAAGGAGCAATAGATTGCTTAATAAGTATAATTCAATGACGATAAATATTTATTGAAAGGGGAAA
 TTCATTATAAAAATGGTATTATTAAATCTTATGAAATTATTGAAAGAGTAAATATTCAAAAGCAT
 TGTGTTTGATCTAATTAGGCTTGCTATAGAATTAAATATTAAAGAAGAGGTTTAGAGAACGCTAGATATTGTT
 GAAAAGAATGTTGATTCTGAGAGCATTATCTGAGATCTTGAATTCTTAGTAACAAGGGAGAGCATGAGT
 TTGCTTAAATTAGCTCTTACTTTCTAAGTATAATTCAAGCTTCAGACAAATATACTTATTGTT
 GGGAAAACTTATGAGTCTGAGAGCAAGCATAAAGATTAAAGGCTTGCAATTACTATAAAATTGTTATTGAT
 AATTACCCCTTGTAGTTATTATGAGAGAGCCAAGATAAGATATTATTAAAGCGTTTTAGT

f929.aa

MTKVLVSAIALLSKDKELIPFYKFLFLFFFFTLLACSKVSKDFIVFNKDVKTSRIDLNPNSNVLEVNMEDFFGD
 IIDLKGYKILSVQQENLNLDVYFEQVVLQAQNFSNLNAYLFIIGFDPKIKAGTILFKTQIDIDPKNSYNMYLEDITG
 DYDFNIVIQGFLDKSVLYVFQKSVLNDVSSYRPPIFFDKVNGTVLINKYARSSAYEENRSRESYPISLEKYEKVGE
 DLIISKIEKYEYSNVQGRYCLSSVSEKVGKIDNNIYKTLKNLSKDEVYKFLHGVWYDVHDYNKMHVKDIDEVLFLS
 FERQSSEINLFRKNSQEVAKIEYISKPAYNTLNVSAKSLFSDLIVYNFWIKIVDKENIEIKIDTSTNSYDNGFSG
 TFKRFDENVLNVKKGSSDIYFIPSGNYVYKDKIYDFSYPHLTYIDENKIYYGIFNIFPLKNNFVLEYEIDMGSYKL
 VESFFLEHSERIVQKQKFSTIILNPIKILKDDVSLVKQKLKLERIEKI

t929.aa

KDKELIPFYKFLFLFFFFTLLACSKVSKDFIVFNKDVKTSRIDLNPNSNVLEVNMEDFFGDIDIDLKGYKILSVQQ
 ENLNLDVYFEQVVLQAQNFSNLNAYLFIIGFDPKIKAGTILFKTQIDIDPKNSYNMYLEDITGDFNIVIQGFLKD
 KSVLYVFQKSVLNDVSSYRPPIFFDKVNGTVLINKYARSSAYEENRSRESYPISLEKYEKVGEDLIISKIEKYEYSN
 VQGRYCLSSVSEKVGKIDNNIYKTLKNLSKDEVYKFLHGVWYDVHDYNKMHVKDIDEVLFLSFERQSSEINLFRKN
 SQEVAKIEYISKPAYNTLNVSAKSLFSDLIVYNFWIKIVDKENIEIKIDTSTNSYDNGFSGTFKRFDENVLNVKK
 GSSDIYFIPSGNYVYKDKIYDFSYPHLTYIDENKIYYGIFNIFPLKNNFVLEYEIDMGSYKLVESFFLEHSERIVQ
 KQKFSTIILNPIKILKDDVSLVKQKLKLERIEKI

f929.nt

TABLE 1. Nucleotide and Amino Acid Sequences

ATGACAAAGGTTGGTTAGTGGATTGCTCTGAGTAAGGATAAAGAATTAACTCCATTATAAATTT
 TGTTTTATCTTTTTTACATTACTTGCTTCCAAGGTAAAGCAAGGATTTATTGTTTAAACAAAGATGT
 AAAGACTTCTCCAGGATCGATAATCCAATTCCAATGTTAGAAGTTAAAGGAAAGATTTGGAGAT
 ATTATAGATTTAAAGGTATAAAATTCTTCAGTTCAAGGAAATTAAATTAGATGTATTTGAGCAGG
 TGGTTTAGCTCAAATTCTAATGCATATTGTTATTGTTGATCCTAAATTAAAGCTGG
 AACGATTCTTTAAACTCAAATAGATATTGATCCAAAATCTTATAACATGTATCTGAAGATATTACAGGT
 GATTATGATTTAATATAGTTCAAGGATTTAAAGATAAATCTGTTGTATGTTTCAAAATCTGTT
 TAAATGATGTCTCTTATAGGCCTATATTGACAAAGTTAATGGAACGTCTTATTAAATAAGTATGCAAG
 ATCTTCAGCTTATGAAGAAAACAGATCAAGAGAAAGCTATCCTATTCTAGAAAATATGAAAAGTGGGGAA
 GATTTAATAATTAGCAAGATTGAAGAAAATATGAATATTCTAATGTCAGGGTAGATATTGTCCTCTGTGAGCG
 AAAAGTTGGTAAATTGATAATAATTATAAAACTTTAAAGAATTAAAGCAAGATGAAGTTATAAATT
 GCATGGAGTTGGTATGATGTTATGACTATAATAAAATGCATGTCAAAGATATTGATGAAGTTTATTCTGTCT
 TTTGAAAGCAATCAAGCAGATTAACTTTCAAGGAAAATTCTCAAGAAGTTGCAAAGATTGAATATAATTCAA
 AACCTGTTACAATACTCTTAAATGTTAGTCAAAGTCTCTTCAAGATTGATAGTTATAACTTTGGATCAA
 AATTGAGATAAAAGAAAACATTGAAATCAAATTGACACTAGCACAAATTCTTATGATAATAGGGATTTCGGGT
 ACATTAAAGAGGTTGATGAGAATGTCTTAAATGTTAAAAGGGAGTAGTGTATTTATTCTACTGGAA
 ATTACGTGATAAGGATAAAATTATGATTTCCTTACCCCCATTAACTTATATTGATGAGAATAAAATT
 TGGCATTTTAATATTTCCTTAAAGGAAATTGTTCTGAATATGAGATTGACATGGTAGTTACAAGCTT
 GTTGAATCTTTCTTGAGCATAGCAGAAATTGTTCAAAGCAAAATTCTACAATCATTAAATCCTA
 TAAATTAAAGATGATGTAAGCTTAGTAAAGGGCAAAATTAAAGCTTGAGCGAATAGAAAAATATGA

t929.nt

AAGGATAAAGAATTAACTCCATTATAAATTGTTTATTCTTTTACATTACTTGCTTCCAAGG
 TAAGCAAAGATTTATTGTTTAAACAAAGATGTAAGAAGACTTCTCCAGGATCGATAATCCAATTCAAATGTTT
 AGAAGTTAAATAAAATGGAAGATTTTGAGAGATATTAGATTAAAGGTATAAAATTCTTCAGTTCAAGCAG
 GAAAATTAAATTAGATGTGATTGAGCAGGTGTTAGCTCAAATTCTTCAAATCTTAAATGCATATTG
 TTATTATTGGTTTGATCTAAATTAAAGCTGGAACGATTCTTTAAAACCTCAAATAGATATTGATC
 AAATTCTTAAACATGTATCTGAAGATATTACAGGTGATTGATTTAAATAGTTATTCAAGGATT
 AAATCTGTTGTATGTTCTAAATTCTGTTAAATGATGTCTTATAGGCCTATATTGACAAAG
 TTAATGGAACCTGTTATTAAAGTATGCAAGATCTCAGCTTATGAGAAAACAGATCAAGAGAAAGCTATCC
 TATTCTTAGAAAATATGAAAAGTGGGGAGATTAAATTAGCAAGATTGAGAAAATATGAATATTCTAAT
 GTTCAGGGTAGATATTGTTCTCTGTGAGCGAAAAGTGGTAAATTGATAATAATTATGAACTTAA
 AGAATTAAAGCAAAGATGAAGTTATAAATTGTCATGGAGTTGGTATGATGTTATGACTATAATAAATGCA
 TGTCAAAGATATTGATGAGTTATTCTGTTGAAAGGCAATCAAGCAGATTAACTTTCAAGGAAAAT
 TCTCAAGAAGTTGCAAAGATTGAAATATTCTAAACCTGCTTACAATACTCTTAAATGTTAGTGC
 AAAGTCTCTTT
 TTTCAAGATTGATAGTTATAACTTTGGATCAAATTGAGATAAGAAAACATTGAAATCAAATTGACACTAG
 CACAAATTCTTATGATAATAGTGGATTTCGGGTACATTAAAGAGGTTGATGAGAATGTCTTAAATGTT
 GGGAGTAGTGTATTATTGTTATTCTCTAGTGGAAATTACGTGATAAGGATAAAATTGATTTCTTACCCCC
 ATTAACTTATATTGATGAGATAAAATTATTGAGCTTAAATTGAGCTTAAATTGTTCTTAAAGGAAATTGTT
 TGAATATGAGATTGACATGGTAGTTACAAGCTGTTGAATCTTTCTTGAGCATAGCAGAAAGAATTGTT
 AAGCAAAATTCTACAATCTTAAATCCTATTAAATTAAAAGATGATGTAAGCTTAGTAAAGGGC
 AATTAAAGCTTGAGCGAATAGAAAAATATGA

f933.aa

MNKLLIFVLATFCVSSFAQANDSKNGAFGMSAGEKLLVYETSKQDPIVPFLNLFLGFGIGSFAQGDI
 LGGSLIL
 GFDAVGIGLILAGAYLDIKALDGITKKAQFWTWGKGVMLAGVVTMAVTRLTEIILPFTFANSYNRKLKN
 LVAL
 GGFEPSPDVAMQSSALGFELSFKKSY

t933.aa

TABLE 1. Nucleotide and Amino Acid Sequences

NDSKNGAFGMSAGEKLLVYETSKQDPIVPFLNLFLFGFGIGSFAQGDILOGGSLILGFDAVGIGLILAGAYLDIKAL
DGITKKAQFWGKGVMLAGVVTMAVTRLTEIILPFTFANSYNRKLKNSLNVALGGFEPSFDVAMGQSSALGFEL
SFKKSY

f933.nt

ATGAATAAACTTTAATTTGTTGGCAACCTTGTGTTCTAGCTTGCTCAAGCTAATGATTCTAAAA
ATGGTGCCTTGGGATGAGTGCCTGGAGAAAAACTTTGGTTATGAAACTAGCAAGCAAGATCCTATTGTACCAATT
TTTATTGAACCTTTTGTAGGGTTGGAATAGGCTCCTTGCTCAAGGAGATATTCTGGAGGTTCTCTTATTCTT
GGATTGATGCGTTGGTATAGGGCTTACTTGCGGGGCTTATTGGATATCAAAGCCTGATGGTATTACTA
AAAAAGCTGCTTTCAATGGACTTGGGTAAGGGAGTTATGTTAGCAGGTGTGGTTACTATGGCTGTGACAAGATT
AACAGAAATTATTCTCCATTACATTGCTAATAGTTATAATAGGAAGCTAAAAAATAGCCTAATGTAGCTTTA
GGAGGATTGAAACCTAGTTGATGTTGCAATGGCCAATCCAGTGCTTGGTTGAACGTGCTTTCAAAAAAA
GCTATTAA

t933.nt

AATGATTCTAAAATGGTGCCTTGGGATGAGTGCCTGGAGAAAAACTTTGGTTATGAAACTAGCAAGCAAGATC
CTATTGTACCATTTTATTGAACCTTTTGTAGGGTTGGAATAGGCTCCTTGCTCAAGGAGATATTCTGGAGG
TTCTCTTATTCTGGATTGATGCGGTTGGTATAGGGCTTACTTGCGGGGCTTATTGGATATCAAAGCGCTT
GATGGTATTACTAAAAAGCTGCTTTCAATGGACTTGGGTAAGGGAGTTATGTTAGCAGGTGTGGTTACTATGG
CTGTGACAAGATTAACAGAAATTATTCTCCATTACATTGCTAATAGTTATAATAGGAAGCTAAAAAATAGCCT
TAATGTAGCTTAGGAGGATTGAAACCTAGTTGATGTTGCAATGGCCAATCCAGTGCTTGGTTGAACGT
TCTTCAAAAAAAGCTATTAA

f940.aa

MRKYIFIILIAVLLIGVNIKKIAAAANIDRHTNSTLGINPDSVGIPIFYNDLSKAYPTNLYPGGIGAIKYQYHILNN
LAIGLELRYMFNFDINHSFNILNPDSVGKIFYSVPITFSINYIFDIGELFQIPVFTNIGFSLNTYGDRNNNITNL
RTFDALPTISFGSGILWNFNYKWAFGATASWWMMFEFGNSAKMAHFALVSLSVTVNVNKL

t940.aa

ANIDRHTNSTLGINPDSVGIPIFYNDLSKAYPTNLYPGGIGAIKYQYHILNNLAIGLELRYMFNFDINHSFNILNP
SSVGKIFYSVPITFSINYIFDIGELFQIPVFTNIGFSLNTYGDRNNNITNLRTFDALPTISFGSGILWNFNYKWAF
GATASWWMMFEFGNSAKMAHFALVSLSVTVNVNKL

f940.nt

ATGAGAAAGTATATTTTATAACTAATTGCACTTGCTAATTGGTGTAAACATAAAAAAAATTGCGGCCGCAG
CCAATATTGATAGGCATAACAACTCCACTTTAGGAATAGTTAAGTGTAGGAATCCCTATTTCACACGACTT
ATCAAAAGCTTATCCTACCAATTATATCCAGGAGGTATTGGGCAATAAAATACCACTTACCAATTAAACATT
TTAGCAATTGGACTTGAACTAAGGTATATGTTAACTTGTATTAACCAATTCTTTAATATATTAAATCCAGATT
CAAGTGTAGGTAAAATTGGTATAGCGTGCCTATTACATTTCATAAAATTATATTTGATATAGGAGAATTATT

TABLE 1. Nucleotide and Amino Acid Sequences

TCAAATTCCAGTCTTCACAAATATAGGGTTTCTCTTAATACATATGGAGATAGAAACAACAATTACAAATTAA
 AGAACTTTGATGCACTCCCTACAATCTCTTGGATCTGGAATTATGGAACCTTAACATATAAATGGGCTTTG
 GAGCAACAGCAGTCTGGTGGATGATGTTGAATTGGAAATTCTGCTAAAATGGCACATTGCACTTGTATCATT
 ATCAGTTACAGTGAATGTAATAAATTGTAG

t940.nt

GCCAATATTGATAGGCATACAAACTCCACTTAGGAATAGTTAAGTAGGAAATCCCTATTTCACAACGACT
 TATCAAAGCTTATCCTACCAATTATCCAGGAGGTATTGGGCAATAAAATACCACTTAAACAA
 TTTAGCAATTGGACTTGAACTAAGGTATATGTTAACCTTGATATTAACCATTCTTTAATATATTAAATCCAGAT
 TCAAGTGTAGGTAATTTTATAGCGTGCCTATTACATTTCATAAAATTATATTTGATATAGGAGAATTAT
 TTCAAAATTCCAGTCTTCACAAATATAGGGTTTCTCTTAATACATATGGAGATAGAAACAACAATTACAAATT
 AAAGAACTTTGATGCACTCCCTACAATCTTTGGATCTGGAATTATGGAACCTTAACATATAAATGGGCTTT
 GGAGCAACAGCAGTCTGGTGGATGATGTTGAATTGGAAATTCTGCTAAAATGGCACATTGCACTTGTATCAT
 ATCAGTTACAGTGAATGTAATAAATTGTAG

f943.aa

MKNQFLNSYFQLITTIFLISSITIAAEEITSTLKVPNGFKVEIFLNNTIEKPRGITSQDGNIFIGSGSTFAYFVT
 KNRKIYTIAKTLQKPIGIDYWDNKLYISSVDKIIYVVKNVKEEINKSIKSHKDYTWKMQIFALLPKNNSQMHSGRYI
 KVDSKNNKLIVNIGSQHNVKIPPKKEAVILSINLKTKEEIVAFGVRNSVGFDFHPISNEIYFSDNGQDGLGDNIP
 PDEINVITEYKEHFGFPYVFGKNQKNYGFYNKAPKNTKFIPSIYELPAHVAPLGIHFYRGNNFPKEYINKLFIAEH
 GSWNRSSPVGYKITLDIDSRTARNYKTFLYGFLKHDKSKFGRPVDIITYYDGSILFTDDFGNKIYRVYYEKI

t943.aa

EITSTLKVPNGFKVEIFLNNTIEKPRGITSQDGNIFIGSGSTFAYFVTKNRKIYTIAKTLQKPIGIDYWDNKLYI
 SSVDKIYVVKNVKEEINKSIKSHKDYTWKMQIFALLPKNNSQMHSGRYIKVDSKNNKLIVNIGSQHNVKIPPKKEA
 VILSINLKTKEEIVAFGVRNSVGFDFHPISNEIYFSDNGQDGLGDNIPPDEINVITEYKEHFGFPYVFGKNQKNY
 GFYNKAPKNTKFIPSIYELPAHVAPLGIHFYRGNNFPKEYINKLFIAEHGSWRSSPVGYKITLDIDSRTARN
 YKTFLYGFLKHDKSKFGRPVDIITYYDGSILFTDDFGNKIYRVYYEKI

f943.nt

ATGAAAAATCAATTAAATAGCTATTTCAATTAACTATTACACTATTCTCATCTATAACTATTGCAG
 CAGAAGAAATAACAAGCACACTAAAGTCCTAATGGATTAAAGTCGAATTTCATAAACTACAATTGAAAA
 ACCTAGAGGAATCACAAGCGATCAAGATGGAAATATATTCACTAGGATCTGGAAGCACTTTGCATACTTGTAAACA
 AAAACAGAAAAATTATACCATAGCAAAACCTGCAAAACCTATTGGTATTGATTATTGGATAATAAAACTCT
 ACATATCTCTGCGATAAAATATATGTAGTTAAAATGTAAAGAAGAAATTAAATAAAAGCATAAAATCACATAA
 AGACTATACATGGAAATGCAATTTCGCACTTTGCCAAAAATAATTCTCAAATGCACTCAGGACGTTACATT
 AAAGTAGATTCTAAAAATAACAATTAAATAGTAAATATAGGATCCCAGCACAATGTTAAATTCCCCAAAAAG
 AAGCAGTAATCCTTAGTATTAAATTAAAAACAAAAAGAAGAAATTAGTAGCTTTGGAGTGAGAAACTCAGTTGG
 GTTGATTTCACCCAATTAGCAATGAAATATATTAGCGACAATGGCAAGACGGATTAGGAGACAACATTCCC
 CCAGATGAAATAAACGTAATAACCGAATATAAGAACATTGGATTCCCTATGTGTTGGAAAAATCAAAAAA
 ATTACGGTTTATAACAAAGCACCCAAAAACACTAAGTTATCCCCTATTTACGAACCTCCGGCACATGTAGC
 TCCACTTGAATACACTTTACCGGGAAATAACTTTCCAAAAGAACATAAATAATTATTCATAGCAGAACAC
 GGCTCGTGAACAGATCTTCTCTGTGCGTACAAAATAACAACACTAGACATTGATTCTAAACCAGAACAGCAA

TABLE 1. Nucleotide and Amino Acid Sequences

GAAATTACAAGACTTTTATATGGATTTAAAGCACGACAAATCTAAATTGGACGCCCTGTTGATATAATCAC
ATATTATGACGGTTCAATTCTTTACAGATGACTTGGAAATAAAATACAGAGTTACTACGAAAAGATTAA

t943.nt

GAAATAACAAGCACACTAAAGTCCTAATGGATTAAAGTCGAAATTAAAAAATACAATTGAAAAACCTA
GAGGAATCACAAGCGATCAAGATGGAAATATATTCAAGATCTGGAAAGCACTTTGCATACTTGTAAACAAAAAA
CAGAAAAATTATACCATAGCAAAACCTGCAAAACCTATTGGTATTGATTATGGGATAATAAAACTCTACATA
TCTCTGTGATAAAATATATGTAGTAAAATGTAAGAAGAAATTAAATAAAAGCATAAAATCACATAAAGACT
ATACATGGAAAATGCAAATTTCGACTTTGCCAAAAATAATTCTCAATGCACCTCAGGACGTTACATTAAAGT
AGATTCTAAAATAACAAATTAAAGTAAATAGGATCCCAGCACAATGTTAAAATTCCCCAAAAAGAAGCA
GTAATCCTTAGTATTAAATTAAAAACAAAAAGAAGAAATAGTAGCTTTGGAGTGAGAAACTCAGTTGGTTG
ATTTTCACCCATTAGCAATGAAATATTTAGCGACAATGCCAAGACGGATTGGAGACAAACATTCCCCAGA
TGAAATAAACGTAATAACCGAATATAAGAACATTGGATTCCCTATGTGTTGGAAAAATCAAAAAAATTAC
GGTTTTATAACAAAGCACCCAAAAACACTAAGTTATCCCCTATTTACGAACCTCCGACATGTAGCTCCAC
TTGGAATACACTTTACCGGGAAATAACTTCCAAAAGAACATACATAAAATTATTACATAGCAGAACACGGCTC
GTGGAACAGATCTCCCTGTTGGCTACAAAATAACAAACACTAGACATTGATTCTAAACCAGAACAGCAAGAAAT
TACAAGACTTTTATATGGATTTAAAGCACGACAAATCTAAATTGGACGCCCTGTTGATATAATCACATATT
ATGACGGTTCAATTCTTTACAGATGACTTGGAAATAAAATACAGAGTTACTACGAAAAGATTAA

f952.aa

MNYARFAVLIVLLFFYIWFFIILRMKRTNLFLLEKIQNGAKILDIRSPKEYSKSHYLKSINIPFNNLFAKKDKLGD
FESPIIVYGKSFNKSYEAKKVLKSMGFKNVFVAQTLKDMQPQAKKEVG

t952.aa

RMKRTNLFLLEKIQNGAKILDIRSPKEYSKSHYLKSINIPFNNLFAKKDKLGFESPIIVYGKSFNKSYEAKKVLK
SMGFKNVFVAQTLKDMQPQAKKEVG

f952.nt

ATGAATTATGCAAGATTGCAAGTATTAAATAGTTCTGCTTTTTATATTGGTTTTTATTATCCTTAGGATGA
AAAGAACTAATCTGTTTGTTAGAAAAAATCCAAATGGAGCAAAATTGGATATTGGCTCTCCAAAGAATA
TAGCAAGTCTCATTATTGAAGTCATTAAACATTCTTTAATAATTATTGGCTAAAAGGATAATTAGGTGAT
TTTGAGTCCCATAATTGTTATGGTAAAAGTTTAATAAGTCTTACGAGGCTAAAAGTTAAAAGCATGG
GATTTAAGAATGTGTTGCTGGAACCTTGAAAGACATGCCACAAGCAAAAAAGAAGTTGGTTGA

t952.nt

AGGATGAAAAGAACTAATCTGTTTGTTAGAAAAAATCCAAATGGAGCAAAATTGGATATTGGCTCTCCCA
AAGAATATAGCAAGTCTCATTATTGAAGTCATTAAACATTCTTTAATAATTATTGGCTAAAAGGATAATT
AGGTGATTGAGTCCCATAATTGTTATGGTAAAAGTTTAATAAGTCTTACGAGGCTAAAAGTTAAA
AGCATGGATTTAAGAATGTGTTGCTGGAACCTTGAAAGACATGCCACAAGCAAAAAAGAAGTTGGTTGA

TABLE 1. Nucleotide and Amino Acid Sequences

f378.aa

MIKKFLLFAMLNIFLTNKAHSNEEIIIEISTEIQKEKYIPFLISRGKTQLEDLVKYTLEINPELDKNYVNTVAKTYI
DESLIEGVNYDIAYAQMLLETGALKFNGIVSKEQHNFSGIGATNNLTKGNSFSNITEGIKAHIQHLKAYASKQNIK
SNMVDPRFYLVKRGSAPTIYDLTGKWAQDKLYDKKKKILLELEYYNNANKS

t378.aa

NEEIIIEISTEIQKEKYIPFLISRGKTQLEDLVKYTLEINPELDKNYVNTVAKTYIDESLIEGVNYDIAYAQMLLET
GALKFNGIVSKEQHNFSGIGATNNLTKGNSFSNITEGIKAHIQHLKAYASKQNIKSNMVDPRFYLVKRGSAPTIYD
LTGKWAQDKLYDKKKKILLELEYYNNANKS.

f378.nt

ATGATAAAAAATTCTGCTATTGCAATGCTAACATCTTTAACAAATAAGCTCATAGTAATGAAGAGATAA
TCGAAATAAGTACTGAAATAACAAAAGGAAAATATATTCCCTTTAACAAATAAGCTCATAGTAATGAAGAGATAA
CCTTGTAAAATATACTCTAGAAATAATCCAGAGCTTGACAAAAACTATGTAATAACTGTTGCTAAAACCTATATA
GACGAATCTTGATTGAAGGGGTTAATTATGACATTGCCTATGCTCAAATGTTACTAGAAACAGGAGCTCTAAAT
TCAATGGAATAGTTCAAAAGAACACAATTTCAGGAATAGGCGCTACTAATAATCTTACAAAAGGAAATTC
TTTTCCAATATTACAGAAGGAATTAAAGCTCATATTCAACATTAAAGCTTATGCTTCAAACAAATATCAA
TCAAATATGGTTGATCCTAGATTTACCTGTTAAAGAGGATCTGCTCCAACAATATGATTGACTGGGAAAT
GGGCAAAAGACAAACTTACGACAAAAACTTAAAGGAAATTATTAGAACTATTAGAAATATAATGCAAATAA
AAGCTAA

t378.nt

AATGAAGAGATAATCGAAATAAGTACTGAAATAACAAAAGGAAAATATATTCCCTTTAACAAATAAGTAGAGGAAAAA
CTCAACTAGAAGACCTTGTAAAATATACTCTAGAAATAATCCAGAGCTTGACAAAAACTATGTAATAACTGTTGC
TAAAACCTATATAGACGAATCTTGATTGAAGGGGTTAATTATGACATTGCCTATGCTCAAATGTTACTAGAAACA
GGAGCTCTAAATTCAATGGAATAGTTCAAAAGAACACAATTTCAGGAATAGGCGCTACTAATAATCTTAA
CAAAGGAAATTCTTTCAATATTACAGAAGGAATTAAAGCTCATATTCAACATTAAAGCTTATGCTTCAA
ACAAAATATCAAATATGGTTGATCCTAGATTTACCTGTTAAAGAGGATCTGCTCCAACAATATGAT
TTGACTGGGAAATGGGAAAAGACAAACTTACGACAAAAACTTAAAGGAAATTATTAGAACTATTAGAAATATA
ATAATGCAAATAAAAGCTAA

f4.aa

MKLFRRNVMIKMPSSFTIIFSLIVFTILTYVI PAGKFDKEFKQMGDGSKREII VAGTYQYVDRGSRGFLHPIMTI
LTAMSKGMEHAVEVIVFVLIVGGAYGIIMKTGAIDVGIYFLIKLGHKDLLIPLLMIIFSIGGTVTGMSEETLPF
YFVMIPLIVALGYDSLVGAIIIALGAGVGTMASTVNPATGIAIASIASLQDGFYFRIVLYFVSVLAITYVCVY
ASKIKKDPSKSLVYSQKDEHYQYFVKKDCLSTGDNQAQNALEFTFAHKLVLLFGFMILILIFSIVNLGWWMQEMTM
LYLGVAIIISAFICKLGETEMWDVFVKGSSESLLTAALVIGLARGVMIVCDGGLITDTMLNAATNFLYNLPRPLFIIL
NEIIQIFIGFVVPSSSGHASLTMPIMAPLADFLSIPRASVVIAMQTASGLINLITPTSGVIMAVLGISRLSYGTWF
KFVLPLFMIEFFFISILVIIANIYLSF

t4.aa

TABLE 1. Nucleotide and Amino Acid Sequences

KFDKEFKQMGDKREIIVAGTYQYVDRGSRGFLHPIMTILTAMSKM**E**HAVEVIVFVLIVGGAYGIIMKTGAIDV
 GIYFLIKKLGHKD**K**LLIPLLMFIFSIGGTVTGMSEETLPFYFVMIPLIVALGYDSL**V**GAIIALGAGVGTMASTVN
 PFATGIA**S**AIASISL**S**QDGFYFRIVLYFVSVLAAITYVCVYASKIKKDP**S**KLV**S**QK**D**EHYQYFVK**D**GL**S**TD**G**NA
 QNA**E**FTFAHKLV**L**LLFGFM**L**IL**I**FSIVNL**G**W**W**QM**E**TM**L**YLG**V**AI**I**SAFICKL**G**ETEMWD**A**FVK**G**SE**S**LLTAAL
 VIGLARGVMIV**C**DD**G**L**I**TD**M**LN**A**AT**N**FLY**N**LP**R**PL**I**IL**N**E**I**IQ**I**FIG**V**VP**S**SS**G**HAS**L**TM**P**IMAP**L**AD**F**LS**I**P
 RASVVIAMQTASGLINLITPTSGVIMAVLG**I**RL**S**YGTWFK**V**PL**F**MI**E**FF**I**SILV**I**ANI**I**YLS**F**

f4.nt

ATGAAATTATTTAGGAGAACGTTATGATCAAATGCCAAGTAGTTTACAATAATTCTTTAATTGTATTTG
 TTACCAATTAAACGTATGTGATTCCCTGCCGTAAAGTTGATAAAGAATTAAAGCAATGGGTGATGGATCTAAAG
 GGAAATAATTGTTGCTGGAACTTATCAATATGTAGATCGAGGCTCTAGGGGATTTTACATCTTATTATGACTATT
 TTAACCGCAATGTCAAAGGGGATGGAACATGCAGTTGAAGTTATTGTTTTAATTGTTGGGGTGTCTATG
 GGATTATTATGAAA**A**CTGGAGCAATAGATGTGGAAATTATTTAATCAAGAAGTTGGGGCACAAAGATAAGTT
 GCTTATTCTTGTAAATGTTATTCTCAATTGGTAACTGTAACCGGAATGAGTGAAGAGACCCCTCCTTT
 TATTTGTTATGATTCCCTGTAGTAGCTTGGGTATGATACTGCTTGTGAGCTCTTGTCTATTAGCTT
 CTGGAGTGGGAACTATGGCTACTGTAAATCCATTGCGACAGGAATTGCATCTGCAATAGCTTCTATTAGCTT
 GCAGGATGGATTATTAGAATTGTTCTTATTGTTATGATCAGTATTGGCTGCTATAACCTATGTTGTGTTAT
 GCGTCTAAAATTAAAAAGGATCCCTCAAATCGCTGTGATTCTCAAAAGATGAACATTATCAATATTGTTA
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 TTTATTGATTATGATATTGATTGATATTGATTGCTATTGTTAATCTGGTTGGATGCAAGAAATGACAATG
 TTGATCTGGAGTTGCTATTATATCGGCTTTATTGTAATTAGGTGAAACTGAAATGTGGATGCGTTGTGA
 AAGGTTCTGAAAGTCTGCTACCGCTGCTTGTATTGGACTTGCTAGAGGTGTTATGATAGTATGTGATGATGG
 GTGATTACAGATACTATGTTAAATGCTGCTACTAATTCTTATACAATCTTCAAGACCCCTTTTATCATATTG
 AATGAAATTATTCAAATATTAGGATTGTTGTCATCTCAGGACATGCTAGTCTCACTATGCCAATAA
 TGGCTCTTGGCATTGTCATTCCAAAGAGCTTCAGTTGTTATTGCCATGCGACACTGCATCTGGCTTAT
 TAATTGATAACACCTACCAGGGAGTTAAATGGCTGATTGGGGATATCCAGATTGAGTTATGGTACGTGGTT
 AAGTTGTTTACCAATTATGATTGAGTTTTATCTATTAGTTATTAGCTAACATTATTAAAGTT
 TTTAG

t4.nt

AAGTTGATAAAGAATTAAAGCAAATGGGTGATGGATCTAAAGGGAAATAATTGTTGCTGGAACTTATCAATATG
 TAGATCGAGGCTCTAGGGATTTCACATCTTATTATGACTATTAAACCGCAATGTCAAAGGGGATGGAACATGC
 AGTTGAAGTTATTGTTCTTAAATTGTTGGGGTGTATGGGATTATTATGAAA**A**CTGGAGCAATAGATGTG
 GGAATTATTCTTAAATCAAGAAGTTGGGCACAAAGATAACTGCTTATTCTTGTAAATTGTTATTGTTCAA
 TTGGTGGAACTGTAACCGGAATGAGTGAAGAGACCCCTCCTTTATTGTTATGATTCCCTGATAGTAGCTT
 GGGTTATGATAGTCTGTTGGAGCGGCTATTATTGCTTATTGAGCTGGAGTGGAACTATGGCTTCACTGTAAAT
 CCATTGCGACAGGAATTGCATCTGCAATTAGCTTCTATTAGCTTGCAGGATGGATTATTAGAATTGTTCTT
 ATTGATCAGTATTGGCTCTAAACCTATTGTTGTTATTGCTTAAATTGTTATTGAGTTATTGATATTGATTGATAT
 GCTTGTGATTCTCAAAAGATGAACATTATCAATTGTTGTTAAATTGAGTTATTGATATTGATATTGATATTGAT
 CAGAATTGCTCTGAGTTACTTTGCTCATAAATTAGTTTACTTTATTGAGTTATTGATATTGATATTGATATTGAT
 TTAGCATTGTTAATTGTTGGATGCAAGAAATGACAATTGTTGATCTTGGAGTTGCTATTATCGGCTT
 TATTGTAATTAGGTGAAACTGAAATGTGGATGCGTTGTGAAAGGTCTGAAAGTCTGCTAACCGCTGCTT
 GTTATTGGACTTGCTAGAGGTGTTATGATAGTATGTGATGATGGGTTGATTACAGATACTATGTTAAATGCTA
 CTAATTGTTATACAATTCTCCAAGACCCCTTTTATCATATTGAAATTATTCAAAATTAGGATTGTT
 TGTTCCATTCTCATCAGGACATGCTAGTCTCACTATGCCAATAATTGGCTCTTGGCGATTGTTGCAATTCCA
 AGAGCTTCAGTTGTTATTGCCATGCGACTGCATCTGGCTTATTGATAACACCTACCAGCGGAGTTAA
 TGGCTGATTGGGGATATCCAGATTGAGTTATGGTACGTGGTTAAGTTGTTTACCAATTATGATTGAGTT
 TTTTATCTATTAGTTATTAGCTAACATTATTAAAGTTTTAG

TABLE 1. Nucleotide and Amino Acid Sequences

f43.aa

MKYFYFLFLLIFNVYAQNVNSPALPSPLLPEITENKVERENSSKGGENFSNVGLDGKYVNDTILYGLDSQVTSI
 IKALKKSSDSQYNFSLKKRLEKTFNAELKREILELFISLKYSGGIDTANYILENYESKRYSNALFGLAISYLKEFD
 DKEKLKKTIDILENKEGNVVSIAAYYLGEELSLEYSKNMMEVFEKYSGNDGARREILIALGKMSAVDYQDRIYEI
 SLDNYEGPSIKA AAAIEALSYLASDKVTENADLYLQSNNNLNVKLAIIASLSKDPSSLKSKIELQGFLRDSDDNIRF
 KAINAIKGHRDSSAKDILYKLSDPSLKVRreasAKALIDMDLGNIEIKNIMFDFKIDNNFKISMFSYLLDKDSLK
 ALSIALEIVNKENINRPSNVLRGVASMLAGKKGNFDNFYSKIIDSKNIDLRLALKGAVYNKSSSLDKLKKIKSE
 TNSEYIKMLLKDY

t43.aa

LPSPPLLPEITENKVERENSSKGGENFSNVGLDGKYVNDTILYGLDSQVTSI IKALKKSSDSQYNFSLKKRLEKTF
 NAELKREILELFISLKYSGGIDTANYILENYESKRYSNALFGLAISYLKEFDDKEKLKKTIDILENKEGNVVSIA
 AYYLGELNSLEYSKNMMEVFEKYSGNDGARREILIALGKMSAVDYQDRIYEISLDNYEGPSIKA AAAIEALSYLASD
 KVVTENADLYLQSNNNLNVKLAIIASLSKDPSSLKSKIELQGFLRDSDDNIRFKAINAIKGHRDSSAKDILYKLKS
 DPSLKVRreasAKALIDMDLGNIEIKNIMFDFKIDNNFKISMFSYLLDKDSLKALSTALEIVNKENINRPSNVLRGV
 ASMLAGKKGNFDNFYSKIIDSKNIDLRLALKGAVYNKSSSLDKLKKIKSETNSEYIKMLLKDY

f43.nt

ATGAAATACTTTATTTTTACTTATTTTAATGTGTATGCTAAAATGTTAATTCTCCAGCTCTTC
 CTAGTCCGCCCTTGTGCCGAAATTACAGAAAATAAGCCTGTTGAGAGAGAAAATTCTCTAAGGGAGAGAATT
 TTCTAATGTTGGTTAGATGGTAAGTATGTTAACGATAACATTCTTATGGGCTTGATAGTCAGTGACAAGCATT
 ATAAAAGCTCTAAAAAAATCAAGCGATAGTCATATAATTCTCTTAAAAAAAGACTTGAGAGAAAATTCTTAAATG
 CTGAGCTAAAAGGAAATACTGAATTGTTATTCTCTTAAAGTATTGGGGGCATTGATAACAGCAAATTATAT
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 GATAAAAGAAAAATTAAAAAAACTCTTATGACATTCTGAAATAAGAGGGCAATGTGGTATCTATTGAGCTT
 ATTATTAGGAGAGCTTAACTCTCTGAGTATTCTTAAACATGATGGAAGTTTGAAAATTCTGGAAATG
 TCGGGCTAGAAGAGAAATACTTATTGCTCTGGAAAAATGTCGCTGTTGATTATCAGGATAGAATTATGAAATT
 TCGCTAGATAATTACGAGGGCCATCAATTAAAGGCTGCTGCAATCGAAGCGTTGTCATATCTGCTTCAGATAAAG
 TAACAGCTTAAAGTGTATCTCAGAGTAATAACAATAATTAAATGTTAAATTAGCTATTATGCTTCTTT
 GTCCAAAGATCCTCTTAAAGCTAAAGAGATTTCAGGATTTTAAGAGATTCTGATGATAATTAGGTT
 AAAGCTTAAATGCAATCAAAGACATAGGGACTCTCTGCAAAGGATATTGATTATAAGCTTAAAGCGATC
 CATCTCTTAAAGTGTGAGGCTCTGCTAAGGCTTAAATTGATATGGATCTGGAAATTGAGATAAAAACAT
 TATGTTGATTAAAGATTGACAATAATTAAATTCATGTTAGTTACCTTTAGATAAGGATTCTCTAAAA
 GCATTGTCATTGCTTAGAAATTGTTAAAGAGAAAATTAAATAGACCCCTCAAATGTTTAAGGGCGTTGCTT
 CAATGTTGGCTGGTAAAAGGTAATTGATAATTCTTATTCTAAAATCATTGACAGCAAAATTGATTAAAG
 GCATTAGCATTAAAGAGCTGTTATAATAAAATCTCATCGCTTCTGATAAGCTTAAAAAAATTAAAAGTGA
 ACGAACTCCGAATATAAAATGCTTTAAAAGATTATTGA

t43.nt

CTTCCTAGTCCGCCCTTGTGCCGAAATTACAGAAAATAAGCCTGTTGAGAGAGAAAATTCTCTAAGGGAGAGA
 ATTTTTCTAATGTTGGTTAGATGGTAAGTATGTTAACGATAACATTCTTATGGGCTTGATAGTCAGTGACAAG
 CATTATAAAAGCTCTTAAAGGAAATCAAGCGATAGTCATATAATTCTCTTAAAGACTTGAGAGAAAATT
 AATGCTGAGCTTAAAGGAAATACTGTTAATTGTTATTCTCTTAAGTATTGAGGAGCTTAAAGGAGTT
 ATATTCTGAAAATTATGAGAGTAAAGATATTCAAACGCTTATTGGCTTGGCAATTGCTATCTTAAGGAGTT

TABLE 1. Nucleotide and Amino Acid Sequences

TGATGATAAAGAAAAATTAAAAAAACTCTTATTGACATTCTGAAAATAAGAGGGCAATGTGGTATCTATTGCA
 GCTTATTATTAGGAGAGCTTAATTCTCTGAGTATTCTAAAACATGATGGAAGTTTGAAAATACTCTGGAA
 ATGATGGGCTAGAAGAGAAATACTTATTGCTCTGAAAAATGTCGCTGTTGATTATCAGGATAGAATTATGAA
 AATTTCGCTAGATAATTACGAGGGCCATCAATTAGGCTGCTGCAATCGAACGCGTGTATCTTCAGAT
 AAAGTAACTGAAAATGCTGATTGTATCTCAGAGTAATAACAATAATTAAATGTTAAATTAGCTATTATTGCTT
 CTTTGTCCAAAGATCCTCTTTAAAGTCTAAAGAGATTACAAGGATTTAAGAGATTCTGATGATAATTAG
 GTTTAAAGCTATTAGCAATCAAAGGACATAGGGACTCTCTGCAAAGGATATTGATTTAAGCTTAAAGC
 GATCCATCTCTAAAGTTAGGGAGGCTCTGCTAAGGCCTTAATTGATATGGATCTGGAAATTGAGATAAAAA
 ACATTATGTTGATTTAAGATTGACAATAATTAAATTCAATGTTAGTTACCTTTAGATAAGGATTCTCT
 AAAAGCATTGTCAATTGCTTAGAAATTGTTAATAAAAGAAAATTAAATAGACCTCAAATGTTAAGGGCGTT
 GCTTCATGTTGGCTGGTAAAAGGTAATTGATAATTATTCTAAATCATTGACAGCAAAATTGATT
 TAAGGCATTAGCATTAAAGGAGCTGTTATAATAATCTCATCGCTTCTGATAAGCTTAAAGGTTAAAG
 TGAAACGAACCTCGAATATTAAATGCTTTAAAGATTATTGA

f50.aa

MKFVNNLFKGCLICFFLFFSCLTDRSIQDSHISDIVEKKKEAVIIDNNNVVLGSNEGFKRDYLIGLKDNESFF
 LSDAFLKENNFYFKKARESYAKKNIGLNTYYLNKIVTNENQHSRELLAKANLFFGYVNYENGFYDLSEYNFDLFLK
 DYKYSHASLRLAELKYLVEKSDAISAFKEINEFSISGYDREIYGFLSNKLGVSHLNLESLGFLDNSVFDTFVFD
 NIFVTNILGGLLRYNIKKNDRCVYLDKDKKSIFLNGIRGFADYNGTIYIGGKNVVYIDDVDGDLKQINVPGNADFS
 NVQVLLAVKNGIFVGTLSNGLWFYDLKNWKNIPLGSNKISSLCFDLSKNNLLVGTVDKAIYSVNVNDNLKIEHLD
 FSKNDNEKNINFIKEYKDSYFVGTYGGGLFELNLNKNSYKKHVIANNIDVNYFMDMEIKDKLLFATFDHGLLIYD
 SENDNWDYFGPNNGLLNLNIKVSRFENYVILGTINNGLVFVDENIKKQL

t50.aa

CLTTDRSIQDSHISDIVEKKKEAVIIDNNNVVLGSNEGFKRDYLIGLKDNESFFLSDAFLKENNFYFKKARESYA
 KKNIGLNTYYLNKIVTNENQHSRELLAKANLFFGYVNYENGFYDLSEYNFDLFLKDYKYSHASLRLAELKYLVEK
 SDAISAFKEINEFSISGYDREIYGFLSNKLGVSHLNLESLGFLDNSVFDTFVFDNIFVTNILGGLLRYNIKKND
 RVYLDKDKKSIFLNGIRGFADYNGTIYIGGKNVVYIDDVDGDLKQINVPGNADFSNVQVLLAVKNGIFVGTLSNGL
 WFYDLKNWKNIPLGSNKISSLCFDLSKNNLLVGTVDKAIYSVNVNDNLKIEHLDFFSKNDNEKNINFIKEYKDSYF
 VGTYGGGLFELNLNKNSYKKHVIANNIDVNYFMDMEIKDKLLFATFDHGLLIYDSENWWDYFGPNNGLLNLNI
 KVSRFENYVILGTINNGLVFVDENIKKQL

f50.nt

ATGAAATTGTTGAATAATTATTAAAGGTTGCTTATATGTTTTCTGTTTTCTGCCTTACTACAG
 ATAGATCTATTCAAGATTCTCATATTAGTATATTGAGAAGAAAAAGAACGAGTCATTATTGATGATAATAA
 TGTTGTTCTGGAGTAATGAGGTAATTAAAGAGACTATTGATAGGATTAAAGATAATGAATCTTTTT
 CTTAGTGTGCTTTAAAGAAAATAATTATTAAAGCCAGGGAAAGTTATGCTAAAAAAATTATG
 GCTTGACAAATTATTATTGAATAAAATAGTAACTAATGAGAATCAGCACAGCAGAGAATTGCTAGCTAACGAA
 TTGTTTTGGATATGAAATTATGAGAATGGTTTATGATCTTCCGAATATAATTGATCTATTAAAG
 GACTATAAATATTCTCATGCTAGTTAAGATTAGCTGAATTAAATATCTGTTAAAGAAAATCTGATGCAATT
 CTGCATTAAAGAGATTATGAATTCTATCTCAGGTTATGATAGAGAGATTATGGCTTTAAGTAATAAAACT
 TGGAGTAAGTCATTAAACTTAGAGTCTTAGGATTCTGACAACAGCGTTTGATACATTGCTTTAATGAC
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 AGGATAAAAAAGCATTAAATGGCATTAGGGGTTTGCAGGATTATAATGGAACAATTATATTGGGTTAA
 AAATGTTGTTATTATAGATGATGTTGATGGGGATTAAAGCAAATAATGTTCCCGTAATGCTGATTAGC
 AATGTCAGTTGCTGTTAAAGGAATTATTGTTGCGACTCTAAATTCTGGATTATGGTTATGATT
 TAAAAAATTGGAAAATACCGCTTGGATCTAATAAAATTCTTCACTCTGCTTGTAGTTAAAAAATTATT

TABLE 1. Nucleotide and Amino Acid Sequences

ATTAGTTGAAACAGTTGACAAGGCTATTATAGTGTAAATGTCGATAATTGAAAAAGATTGAACATTGGATTT
 TTTAGCAAAAATGATAATGAAAAAAATTTAATTAAAGAATATAAGATAGTTATTTGTTGAAACATATG
 GTGGGGCTTTGAAATTAAATTTAAATAAGTACAAAAGCACGTTATTGCCAATAATATTGATGTTAA
 TTATTTATGGATATGGAGATTAAGATAAAAGCTATTGTTGCAACCTTGATCATGGTTATTGATTATGAT
 TCTGAAAATGACAACGGATTATTTGACCAATAATGGCTTCTTAATTGAAATTAAATAAGTTCTAGAT
 TTGAAAATTATGTCATACTGGCACTATTAAACGGTTGGTTGTAGATGAAAATATTAAAAACAGTTATG
 A

t50.nt

TGCCTTACTACAGATAGATCTATTCAAGATTCTCATATTAGTGTAGAGAAGAAAAAGAAGCAGTCATT
 TTGATGATAATAATGTTGTTCTGGGAGTAATGAGGGTAAATTAAAGAGACTATTGATAGGATTAAGATAA
 TGAATCTTTCTTAGTGTGCTTTAAAAGAAAATAATTAAAGCCAGGGAAAGTTATGCT
 AAAAAAAATATTGGCTTGACAAATTATTGAATAAAATAGTAACTAATGAGAATCAGCACAGCAGAGAATTGC
 TAGCTAAAGCGAATTGTTTGGATATGTAATTGAGAATGGTTTATGATCTTCCGAATATAATTGTA
 TCTATTAAAGACTATAAAATTCTCATGCTAGTTAAGATTAGCTGAATTAAATATCTGTTAAAGAAAAA
 TCTGATGCAATTCTGCATTAAAGAGATTAATGAATTCTCATCTCAGTTATGATAGAGAGATTATGGCTTT
 TAAGTAATAAAACTGGAGTAAGTCATTAAACTTAGCTTTAGGATTCTGACACAGCGTTTGTACATT
 TGTCTTAATGACAATATATTGTAACTAATATATTGGAGGGCTTTAAGATATAATTAAAAAAATGATTGT
 AGAGTCTATCTAAGGATAAAAAAGCATTTAAATGGCATTAGGGTTTGGGATTATAATGAAACAATT
 ATATTGGTGGTAAAAATGTTTATTATAGATGATGTTGATGGGATTAAAGCAAAATAATGTTCCCGTAA
 TGCTGATTTAGCAATGTACAAGTTGCTTGCTGTTAAAATGGAATATTGTTGGCACTCTAAATTCTGGATT
 TGGTTTATGATTAAAAATGGAAAAATACCGCTTGGATCTAATAAAATTCTCACTCTGCTTGATAGTT
 TAAAAAATTATTAGTTGAAACAGTTGACAAGGCTATTATAGTGTAAATGTCATAATTGAAAAGATTGA
 ACATTGGATTTTAGCAAAATGATAATGAAAAAAATATTAAATTAAAGAATATAAGATAGTTATT
 GTTGGAACATATGGGGGCTTTGAATTAAATTAAAGATACAAAAGCACGTTATTGCCAATA
 ATATTGATGTTAATTATTGATATGGAGATTAAGATAAAAGCTATTGTTGCAACCTTGATCATGGTT
 ATTGATTATGATTCTGAAAATGACAACGGATTATTTGACCAATAATGGCTTCTTAATTGAAATTAAATA
 AAAGTTCTAGATTGAAAATTATGTCATACTGGCACTATTAAACGGTTGGTTGTAGATGAAAATATT
 AAAACAGTTATGA

f65.aa

MIFKNVPFQINLILFLVSKVAKINASSKFYYAEQWYVIFNSQMKKPENYKKNIFFLQKALKYPFGNPKYSLTKI
 ETKEQWEKYKLLFKMHVNLLLVRQNLHLDLFDTNLYFFKTPKDGIISNLEKSKKLYKLAINYYSEALKYHKKL
 ENYTTVKLENDGITNWEDEYHKISLKELNYDIIKKELLRIDETKAFFEQQGPNEY

t65.aa

KINASSKFYYAEQWYVIFNSQMKKPENYKKNIFFLQKALKYPFGNPKYSLTKIETKEQWEKYKLLFKMHVNLLL
 RQNLHLDLFDTNLYFFKTPKDGIISNLEKSKKLYKLAINYYSEALKYHKKLENYYTTVKLENDGITNWEDEYHK
 ISLKELNYDIIKKELLRIDETKAFFEQQGPNEY

f65.nt

ATGCATATTTCAAAAATGCCCCCTTCCAAATAATTAAATTATTTATTTCTTTAGTATCAGTTGCAAGATAATG
 CATCGTCAAATTATTACGAGAACATGGTATGTAATTAAATTCTCAAATGAAAAAAACCTGAAACTA
 TAAAAAAATATTCTTCTTCAAAAGCCTTAAACCCATTGGAAATCCAAAATATTCTCTAACTAAAATA
 GAAACCAAGAACAGTGGAAAAATATAAACTCTTCAAAATGCTAAACTTGCTCTAGTTAGGCAAAATT
 TACATTAGGAGATTATTGACACAAAGAAATTATTTCTTCAAAACTCCAGAAAAGATGGAATTATTCCAA
 TCTAGAAAATCAAAAAATTATATAAACTAGCTATTAAATTACTACAGCGAAGCACTAAAATACCACAAAAACTT
 GAAAATTACACAACTGTTAAACTAGAAAAGATGGAATAACAAACTGGGAAGATGAATATCATAAAATTCTCTTA

TABLE 1. Nucleotide and Amino Acid Sequences

AAGAGCTTAATTACTATGACATTATTAAAAAGAACTACTAAGAATTGACGAAACTAAAGCATTGGAAACAAGG
GCCAAACTATTATAA

t65.nt

KINASSKFYYAEQWYVIFNSQMKKKPENYKKNIFFLQKALKYPFGNPKYSLTKIEKTEQWEKYKLLFKMHVNLLLV
RQNLLHGDLFDTRNLYFFKTPKDGIISNLEKSKKLYKLAINYYSEALKYHKKLENYTTVKLENDGITNWEDEYHK
ISLKELNYYDIKKELLRIDETKAFFEQQPNYY

f8.aa

MKNINRLILLIILTHTLLFSCALIADNKSKNLSTSEIILTQKTLLESSLIKNPSNVEYRIPISSIQEILNNNNDSF
LIKTKAAKIKISPQKLEEIKNYLNAYKNYLNNETEWIKFIDQSSVNGNLTIKIDTAFEEKTNFNHTNSDNENLTEL
IELQMHLKEILNLIEQTFHDKNLGYIQLSHINSFPQENINSITKEIIDGKEYIAPHIIANQLLKIKDKKYFEQF
MHFLKVENSKIKTIIIEKQKISDLHNELYYSKQSPPRRRKSTADSDNNNKYDIIPKIIDPNTGIEITPKNLRISILS
NGDIILIKPKIDWTEFFYFWQHVGIFDEEKYEATKKIAFNGIDSFDIKSIIITSNQIKFDTASTQGSGYEKLSTYVQ
SRILKIFSPITDIRTIQKAIINFGRSRYIDNNFGYMVPLISSNLWTDNFNLEEHNKTYCSLMVDRIYKIAGLNVR
NYEISGIITPGEINAAAYNFMSYTIAGILPSVLPKRLIKPTLKEKFIGYNKEIVDAIELKKSKKEKIFGRACNITN
LWCSGS

t8.aa

CALIADNKSKNLSTSEIILTQKTLLESSLIKNPSNVEYRIPISSIQEILNNNNDSFLIKTKAAKIKISPQKLEEIK
NYLNAYKNYLNNETEWIKFIDQSSVNGNLTIKIDTAFEEKTNFNHTNSDNENLTELIELQMHLKEILNLIEQTFH
DKNLGYIQLSHINSFPQENINSITKEIIDGKEYIAPHIIANQLLKIKDKKYFEQFMHFLKVENSKIKTIIIEKQK
SDLHNELYYSKQSPPRRRKSTADSDNNNKYDIIPKIIDPNTGIEITPKNLRISILSNGDIILIKPKIDWTEFFYFW
QHVGIFDEEKYEATKKIAFNGIDSFDIKSIIITSNQIKFDTASTQGSGYEKLSTYVQSRILKIFSPITDIRTIQKAI
NFGRSRYIDNNFGYMVPLISSNLWTDNFNLEEHNKTYCSLMVDRIYKIAGLNVRNYEISGIITPGEINAAAYNF
YMSYTIAGILPSVLPKRLIKPTLKEKFIGYNKEIVDAIELKKSKKEKIFGRACNITNLWCSGS

f8.nt

ATGAAGAATATTAATAGATTAATATTAACTACACACACTTTATTATTCTCTTGTGCCCTAATTGCAG
ATAATAAGCAAAAAATTAAAGCACATCAGAAATCATATTAAACACAAAAACACTACTAGAAAGCTCTTAATAAAA
AAATCCTTCTAATGTAGAATATCGAATACCAATATCCAGTATCCAAGAAATTAAACATAACAATGATTCTTT
TTAATAAAAAAACAGCAGCAAAATCAAATAAGCCCTCAAAACTTGAAGAAATAAAAAACTATCTAAATGCTT
ATAAAAATTATCTAAATAATGAACAGAACGAAATGGATAAAGTTATAGATCAAAGTAGCGCTAACGGAAATTAAACAA
TAAAATTGATACTGCTTTGAAAAAAACAAATTAAATCATACAAATTCAAGATAATGAAAATTAAACAGAACTA
ATAGAACTACAAATGCATCTGGAAAAAGAAATTAAACTTAATTGAGCAAACATTCTGATAAAAATTAGGAT
ATATACAATTAAAGTCACATCAACTCATTCTTCCTCAAGAAAATATAAACTCAATAACAAAAGAAATAATAGATGG
AAAAGAATATATTGCACCGCACATAATAGCAAATCAATTATTTAAAGATAAAAATATTGACAATT
ATGCACCTTTAAAGTTGAAAACAGCAAAATAAAACAAATAATTGAAAAACAAAAATTTCAGATCTTCACAATG
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TCAAGAATATTAAAATATTCTCACCAATAACAGACATAAGAACAAATTCAAAAAGCTATTAAATTGGAAAGAAGTA
GATACATTGACAATAACTTGGATATATGGTTCCATTAAATACCTCTAATTATGGACAGATTCATTCAATCTGAA
AGAAATTCAACAAAACCTATTGCTCTTAATGGTTGATAGAATATAAAATAGCAGGACTTAATGTATCAAGA

TABLE 1. Nucleotide and Amino Acid Sequences

AATTACGAAATTCTGGGAATAATTACTCCTGGAGAAATAATGCAGCAGCTTACAATTACATGTCTTACAGA
 TTGCAGGAATACTTCAAGCGTCTCCAAAAGGCTCATTAACCAACATTAAAGAAAATTCAATTGGTTACAA
 TAAAGAAAATAGTAGATGCAATAGAATTAAAAAAATCGAAAGAAAAATTGGGAGAGCTTGCACATTACAAAT
 CTCTGGTGCTCAGGAAGTTAA

t8.nt

TGTGCCTTAATTGCAGATAATAAGTCAAAAATTAAAGCACATCAGAAATCATATTAAACACAAAAACACTACTAG
 AAAGCTCTTAATAAAAAATCCTCTAATGTAGAATATCGAATACCAATATCCAGTATCCAAGAAATTAAACAA
 TAACAATGATTCTTTAATAAAAAAAACAGCAGCAAAATCAAATAAGCCCTCAAAACTTGAAGAAATAAAA
 AACTATCTAAATGCTTATAAAATTCTAAATAATGAAACAGAAATGGATAAAGTTATAGATCAAAGTAGCGTCA
 ATGGAAATTAAACAATAAAATTGATACTGCTTTGAAAAAAACAAATTAAATCATACAAATTCAAGATAATGA
 AAATTAAACAGAACTAATAGAACTACAAATGCATCTGAAAAAGAAATTAAACTTAAATTGACCAAACATTTCAT
 GATAAAAATTAGGATATATACAATTAAAGTCACATCAACTCATCTTCCTCAAGAAAATATAACTCAATAACAA
 AAGAAATAATAGATGAAAAGAATATATTGACCCGACATAATAGCAAATCAATTATAAAAATAAGATAAAA
 ATATTGACAATTATGCACTTTAAAGTTGAAAACAGCAAAATAAAAACAATAATTGAAAACAAAAATT
 TCAGATCTCACAATGAACTGTATTATTCAAAACAATCCCGCCAGAAGAAGAAAAGGTCACTGCCGATTCCG
 ATAATAACAATAATACGATATAATACCAAAAATAATAGACCCAAATACAGGCATTGAAATAACTCCTAAAATT
 AAGATCTATTATCAATGGCAGACATAATACTAATAAAACAAAATAGATTGGACAGAATTTTTATTGG
 CAACATGTGGAATATTGATGAAGAAAATATGACCACTAAAAAATGCCATTCAATGGAATTGATAGCTTG
 ATATAAAATCAATAATTACAAGCAATCAAATCGATACAGCATCTACTCAAGGTTCAAGGATACGAAAAGCT
 TTCAACATACGTACAATCAAGAATATTAAAAATATTCTCACCAATAACAGACATAAGAACAAATTCAAAAGCTATT
 AATTTGGAAGAAGTAGATACTTGACAAACTTGGATATGGTTCCATTAAATCCTCTAATTATGGACAG
 ATTCAATTCAATCTGAAAGAAATTCAACAAAACCTATTGCTTTAATGGTTGATAGAATATATAAAATAGCAGG
 ACTTAATGTATCAAGAAATTACGAAATTTCGGGAAATAATTACTCTGGAGAAATAATGCAGCAGCTTACAATT
 TACATGTCTTACGATTGCAAGGAAACTTCCAAGCGTCTCAAAAAGGCTTAAACCAACATTAAAAGAAA
 AATTCAATTGGTTACAATAAAAGAAATAGTAGATGCAATAGAATTAAAAAAATCGAAAGAAAAATTGGGAGAGC
 TTGCAACATTACAAATCTCTGGTGCTCAGGAAGTTAA

f82.aa

MTRVFSKFFLFFCFSMLFANSEDSNEKDIVSKDENPVFENEVLGYWVGYNDVSNIKNSIIYIYKYNGEVYGRILT
 IIKDGGKYDAKNPSGDTVVGFESENLAIEGLDFMWGLKYSSSSKKWDRGKIIDPKNGKIIYNSEMRVDSKTGNLITKGK
 VWIFGRSKIWTRAKDDEIPKLDLHNLPAPPVKK

t82.aa

EDSNEKDIVSKDENPVFENEVLGYWVGYNDVSNIKNSIIYIYKYNGEVYGRILTIIKDGGKYDAKNPSGDTVVGFE
 NLAIEGLDFMWGLKYSSSSKKWDRGKIIDPKNGKIIYNSEMRVDSKTGNLITKGKVVIFGRSKIWTRAKDDEIPKLD
 LHNLPAPPVKK

f82nt

ATGACTAGAGTTTTCAAAGTTTTCTTTTTGTTCAATGCTTTATTGCAAATTCAAGAAGATTCAA
 ATGAAAAGGACATTGTTAGCAAGGATGAAAACCCCTGTTTGAAAATGAAGTTAGGATATTGGGTTGGTTATAA
 TGATGTAAGTAACATAAGAATTCTATTATCTATTTATAAATATGGGAAGTTATGGCCGAATTAACT
 ATAATAAAAGATGGCAAAAGTATGATGCTAAAATCCTCAGGAGATACTGTAGTTGGGTTGAAAATCTTGCAA
 TAGAGGGTCTGATTATGTGGGTCTTAAGTATTCTCTTCTAAAGTGGGATAGGGCAAAATAATAGA

TABLE 1. Nucleotide and Amino Acid Sequences

TCCTAAAAACGGTAAAATTATAATTCTGAGATGCGTGTGATAGTAAAACCGGAAATCTTATTACCAAGGGGAAA
 GTTTGGATTGGTAGAAGTAAAATTGGACAAGAGCTAAAGATGATGAAATACCAAAATTAGATTGCATAATC
 TTGTTCCAGCGCCCCCTGTGAAAAATAA

f82.nt

GAAGATTCAAATGAAAAGGACATTGTTAGCAAGGATGAAAACCTGTTTGAAGTTAGGATATTGGG
 TTGGTTATAATGATGTAAGTAAACATAAAGAATTCTATTCTATATTATAAATATAATGGGGAGTTATGGCCG
 AATTTAACTATAATAAAAGATGGCAAAAAGTATGATGCTAAAATCCTCAGGAGACTGTAGTTGGGTTGAA
 AATCTGCAATAGAGGGTCTTGATTTATGTGGGTCTTAAGTATTCTCTTCTAAAAGTGGGATAGGGCA
 AAATAATAGATCCTAAAACGGTAAAATTATAATTCTGAGATGCGTGTGATAGTAAAACCGGAAATCTTATTAC
 CAAGGGGAAAGTTGGATTGGTAGAAGTAAAATTGGACAAGAGCTAAAGATGATGAAATACCAAAATTAGAT
 TTGCATAATCTGTTCCAGCGCCCCCTGTGAAAAATAA

f86.aa

MNKLMMLITFATSLAQTNKASTGLKTDQSFNNSLSESVKLKEIADIYPTNTNFLTGIGIVAGLAGKGDSIKQKD
 LIKILEENNIINEIGSNNIESKNIALVNVLQVKGNTIKGSKHKACVASILDSDKDLTNGILLKTNLKNKEGEIIA
 IASGITQPNNKLKGSGYTIIDSVIINENQNIHNSYNIILKKGNYTLINRIHKILTSKINNKKIKSDSTIEIEAKNIS
 LLEEEIENIKIETNPKILIDKKNGIILASENAKIGTFTFSIEKDQNQNIFLSKNNKTTIQVNSMKLNEFILKNSNNLS
 NKELIQIIQAAQKINKLNGELILEEIDGNQ

t86.aa

LKTDQSFNNSLSESVKLKEIADIYPTNTNFLTGIGIVAGLAGKGDSIKQKDLILIKILEENNIINEIGSNNIESKNI
 ALVNVLQVKGNTIKGSKHKACVASILDSDKDLTNGILLKTNLKNKEGEIIIAASGITQPNNKLKGSGYTIIDSVIIN
 ENQNIHNSYNIILKKGNYTLINRIHKILTSKINNKKIKSDSTIEIEAKNISLLEEEIENIKIETNPKILIDKKNGII
 LASENAKIGTFTFSIEKDQNQNIFLSKNNKTTIQVNSMKLNEFILKNSNNLSNKELIQIIQAAQKINKLNGELILEE
 IDGNQ

f86.nt

ATGAACAAACTAATGTTGATGTTAATTACATTGCAACGAGTCATTAGCCAAACAAACAAAGCTTCAACAGGAC
 TAAAAACAGATCAATCATTAAACATAGCCTATCTGAAAGCGTAAAATTAAAAGAAATTGGGATATTATCCAC
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 CTTATAATTAAATTAGAAGAAAACAATATAATAATGAAATTAGGCTCTAAACATAGAAAGTAAAATATTG
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 ACTGGACTCAAAGATTAAACAAATGAAACTTTAAAACAAATCTTAAAATAAGAGGGGAAATAATAGCA
 ATTGCATCAGGAATTACACAGCCAATAATAATTAAAAGGATCTGGATATACTATAGATAGTGAATAATAATAG
 AGAATCAAATATTAACCACAGTTATAATATAATTCTTAAAAAGGAAATTACATTAATAATAGAATTCTCAA
 AATATTAAACCTCTAAAAATCAACAACAAATTAAATCAGACAGCACAATAGAAATAGAAGCAAAAACATAAGC
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 TAGCAAGTGAACAAATAGGAACCTTTACATTTCATTGAAAAGACAATCAAACATTTTTAAGTAA
 AAATAACAAACAAACAAATTCAAGTAAACTCAATGAAATTAAATGAATTATATTAAAAAAATTCCAACAACTTAC
 AATAAGAATTAAATTCAAAATTCAAGCTGCGAAAAATTAAATAATTAAATGGGAACTTATCTGGAGGAA
 TTGATGGAAACCAAAATTAA

t86.nt

TABLE 1. Nucleotide and Amino Acid Sequences

CTAAAAACAGATCAATCATTAAACAATAGCCTATCTGAAAGCGTAAAATTAAAAGAAATTGCGGATATTTATCCCA
 CAAATACAAATTTTAACAGGTATTGGAATAGTAGCGGGACTGCTGGAAAAGGAGACTCTATAAAACAAAAAGA
 CCTTATAATTAAAATTAGAAGAAAACAATATAAAATGAATAGGCTCAATAAACATAGAAAGTAAAATATT
 GCACTAGTAAATGTCAGTCTCCAAGTAAAAGGTAATACAATCAAAGGTTCAAAACATAAGCTTGCCTGCATCAA
 TACTGGACTCAAAAGATTAAACAAATGGAATACCTTAAACAAATCTTAAAGAGGGGAAATAATAGC
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 GAGAATCAAATATTAAACCACAGTTATAATATAATTCTTAAAGGAAATTATACATTAATAATAGAATTCTATA
 AAATATTAAACCTCTAAAAATCAACAACAAATTAAATCAGACAGCACAAATAGAAATAGAAGCAAAAACATAAG
 CCTATTAGAAGAGATTGAAAATATTAAAGAAACCAACCCAAAGATATTAAATAGACAAAAAAATGGTATTATT
 TTAGCAAGTGAAAATGCAAAATAGGAACCTTACATTTCATTGAAAAGACAATCAAACATTTTTAAGTA
 AAAATAACAAAACAATTCAAGTAACTCAATGAAATTAAATGAATTATATTAAAAAATCCAACAACTTCA
 CAATAAAAGAATTAAATTCAAATAATTCAAGCTGCCAAAAATTAAATAAAATTAAATGGGGAACTTATCTTGAGGAA
 ATTGATGAAACCAAAATTAA

f90.aa

MCPITFTIPFFLAIFFAFSSSFVTDSSVSLLSRNTSLFSTLPISLPIISGTLPAIVTLSKKYLSISLSFSKMFIFI
 KSLFEVIKLPPIWLFIIFASGYFLNAFSIFLCISSLFSFMFI

t90.aa

SSFVTDSSVSLLSRNTSLFSTLPISLPIISGTLPAIVTLSKKYLSISLSFSKMFIFIKSLFEVIKLPPIWLFIIFAS
 GYFLNAFSIFLCISSLFSFMFI

f90.nt

ATGTGTCCTATTACTTTACCATTCCATTTCCTAGCAATATTTTGCTTTCAAGCTCCTTGTACGGACT
 CTTCTGTGCTTTGCTATCAAGAAATACGCTCTTTCTACTTTAACCTCAATTCTTGCTATTATTCTGG
 TACGCTTCCTGCAATAGTTACGCTGTCGAAAAAATATCTGTCAATCTCTTAAGCTTTCTAAATGATTTCATC
 AAATCTTATTGAAAGTATTAAACTTCCATATGGTTATTCAATTATTGTCATCAGGAACTTTAAATGCTT
 TTTCGATTTTGCTATTCTCTTTATCTTATGTTATATGA

t90.nt

AGCTCCTTGTACGGACTCTCTGTCTTGCTATCAAGAAATACGCTCTTTCTACTTTAACCTCAATT
 CTTTGCTATTATTCTGGTACGCTCTGCAATAGTTACGCTGTCGAAAAAATATCTGTCAATCTCTTAAGCTT
 TTCTAAAATGATTTCATCAAATCTTATTGAAAGTATTAAACTTCCATATGGTTATTCAATTATTGTCATCA
 GGATACTTTAAATGCTTTGCTATTCTCTTTATCTTATGTTATATGA

f469.aa

MANVALSSGFISQKIFGIIIMVFLPTIIATPIINFLFKINKSGLKKELPIDQNTHICVSFEYDNLAKILIWDFKN
 ELRKEGFFTQQIKNDSSQYINARKNNISFSIKREGSKITFECPPNNHLIIIQDLFRETILNLEKITKEVETVSLRAK
 KLDYSINYDKILSNINLNKRIKKENIILELKSSNKADVRELLSVNIEIDKERIFQDLMEREKLITTALEKGFAI
 PHLKTNLISKIHIAIGISHEGIDFNALDKNLSHVFILELCPAKDYVSYPRILASVVGKVDLYKKEILNAKTDKEIY
 NIIVSZ

t469.aa

TABLE 1. Nucleotide and Amino Acid Sequences

VFLPTIATPIINFLFKINKSGLKELPIDQNTHICVSFEYDNLAKILIWDFKNELRKEGFFTQQIKNDSSQYINA
 RKNNSFSIKREGSKITFECPNHLIIIQDLFRETIILNEKITKEVETVSLRAKKLDYSINYDKILSNINLNKRIK
 KENIILELKSSNKADVRELLSVNIEIDKERIFQDLMEREKLITTALKEGFAIPHLKTNLISKIHIAIGISHEGI
 DFNALDKNLSHVFLILCPAKDYVSYPRILASVVGKVDLYKKEILNAKTDKEIYNIIVSZ

f469.nt

ATGGCAAATGTAGCATTATCTTCAGGATTATTAGCAAAAAATATTGGAATCATAATAATGGTGTGTTGC
 CAACAATCATTGCAACACCCATAATAAACTTTTATTAAAATAAAAGTGGACTAAAAAGAACTCCCAAT
 AGATCAAAATACACACATATGCGTATCATTGAATATGATAATTAGCCTAAATTCTTATATGGGACTTTAAAAT
 GAGTTAAGAAAAGAAGGATTTTACACAACAAATTAAAATGATTCTTCACAATATATTATGCAAGAAAAACA
 ATATATCCTCTCAATAAAACGAGAAGGTAGCAAAATCACATTGAATGCCAAATAATCATTAAATTATAAC
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 TAAAGAAAGAATATTCAAGATTAAATGAAAGAGAAAAGTTAATTACTACTGCACATAAGAAGGCTTGCCTT
 CCCCATTTAAAACAATTAAATATCAAAATACATATTGAATAGGAATAAGCCATGAGGGAAATTGACTTTAATG
 CTCTTGACAAGAACTTAAGTCATGTTTATATTAAATACTGTGCCAGCAAAGATTACGTTAGCTACCCCTAGAAT
 TTTAGCATCTGTTGGGCAAAGTTGATCTGTACAAAAAGAAATTAAATGCAAAACAGATAAGAAATTAT
 AATATAATAGTGAGCTAA

t469.nt

TTTTGCCAACATCATTGCAACACCCATAATAAACTTTTATTAAAATAAAAGTGGACTTAAAAAGAAC
 TCCCAATAGATCAAAATACACACATATGCGTATCATTGAATATGATAATTAGCCTAAATTCTTATATGGGACTT
 TAAAATGAGTTAAGAAAAGAAGGATTTTACACAACAAATTAAAATGATTCTTCACAATATATTATGCAAGA
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 AGCAAAAAACTAGATTACTCAATAATTACGATAAAATCCTTAGTAATATCAACCTAAATAAAAGAATAAAAAG
 GAAAACATTATTCTAGAATTAAAATCAAGCAATAAGGCTGATGTAATAAGAGAGCTCTAAGCGTAATAAACATTG
 AAATTGATAAAAGAAATATTCAAGATTAAATGAAAGAGAAAAGTTAATTACTACTGCACATAAGAAGGCTT
 TGCCATTCCCCATTAAAACAATTAAATATCAAAATACATATTGAATAGGAATAAGCCATGAGGGAAATTGAC
 TTTAATGCTCTGACAAGAACTTAAGTCATGTTTATATTAAATACTGTGCCAGCAAAGATTACGTTAGCTACC
 CTAGAATTAGCATCTGTTGGGCAAAGTTGATCTGTACAAAAAGAAATTAAATGCAAAACAGATAAGAA
 AATTATAATATAATAGTGAGCTAA

f477.aa

MEKPQGVSIVGAIISGAMHVHLMAEHYGVPVVLHTDHCAKNLLPWVEGLLEYGEKYYSQHKKPLFSSHMLDLSEEP
 KENIEISKKFLERMAKIEMFLEIELGITGGEEDGVNSDRALHELPSTPEDIYYGYSELLKVSBNFQIAAAFGNVH
 GYKPGNVKLTPKVLKDQDYVISKTGVNMAKPVSYVFHGGSGSTIDEINEALSYGVVKMNIDDTQWAWEGLV
 YYKKNESRLQGQLGDKDIDIPNKKFYDPRVWLREAEVSMKDRVKIACKNLNNINRNZ

t477.aa

MHVHLMAEHYGVPVVLHTDHCAKNLLPWVEGLLEYGEKYYSQHKKPLFSSHMLDLSEEP
 KENIEISKKFLERMAKIEMFLEIELGITGGEEDGVNSDRALHELPSTPEDIYYGYSELLKVSBNFQIAAAFGNVH
 DGQDYVISKTGVNMAKPVSYVFHGGSGSTIDEINEALSYGVVKMNIDDTQWAWEGLV
 YYKKNESRLQGQLGDGDIDIPNKKFYDPRVWLREAEVSMKDRVKIACKNLNNINRNZ

f477.nt

ATGGAAAAACCACAAGGAGTTCAATAGTTGGAGCTATTCTGGTGCTATGCATGTTCAATTAAATGGCAGAGCATT
 ATGGTGTCTGTTCTTCATACTGATCACTGTGCTAAAATTGCTCCTGGGTGAAGGCCTTTAGAATA
 TGGAGAGAAATACTATAGTCAGCACAAAAACATTATTCTTCACATATGTTAGATTATCAGAACACCTATT

TABLE 1. Nucleotide and Amino Acid Sequences

AAAGAAAATATTGAAATTCTAAAAAATTCTTAGAAAGAATGGCAAAATTGAAATGTTTGAAATAGAGCTTG
 GAATTACGGGTGGGAAGAGGATGGAGTTGACAATTGAGATAGAGCTTGCATGAACTATTTCTACTCCTGAGGA
 TATTATGGATATTGAGACTTTAAAGTTAGCCAAATTTCAGATTGCAGCAGCTTGGAAATGTTCA
 GGGTATATAAACCGGGAAATGTTAAGCTACTCCAAAGTTAAAGATGGTCAAGATTGTCATATCAAAAA
 CAGGAGTAAATATGGCTAACGCCAGTTCTATGTTTCTATGGAGGGCTGGATCTACATTGATGAGATTATGA
 GGCCTTCTATGGCTGTAAAGATGAATATTGACACAGATACACAGTGGCTGCCTGGAGGGTGTAAAT
 TATTACAAAAAAATGAAAGCTGTTGCAAGGTCAATTAGGAGATGGCAAGGATATTGATATTCAAATAAGAAAT
 TTTATGATCCAAGGGTTGGTAAAGAGAACGCTGAAGTTCTATGAAAGACCGTGTGAAGATTGCATGCAAAATCT
 TAATAATATTAATAGAAATTAA

t477.nt

ATGCATGTCATTAATGGCAGAGCATTATGGTGTCTGTTCTTCATACTGATCACTGTGCTAAAAATTGCT
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 TATGTTAGATTATCAGAAGAACCTATTAAGAAATATTGAAATTCTAGAAAGAATGGCAAA
 ATTGAAATGTTTGAAATAGAGCTTGGATTACGGGTGGGAAGAGGATGGAGTTGACAATTGAGACTT
 TGCATGAACTATTTCTACTCCTGAGGATATTGATATTGAGACTTTAAAGTTAGCCAAATTTC
 GATTGCAGCAGCTTGGAAATGTTCATGGGTATATAACCGGGATGTTAAGCTACTCCAAAGTTAAAG
 GATGGTCAAGATTATGTCATATCAAAACAGGAGTAATGGCTAAGCAGTTCTATGTTTCTATGGAGGGT
 CTGGATCTACAATTGATGAGATAATGAGGCGTTCTATGGCTGTAAAGATGAATATTGACACAGATACACA
 GTGGGCTGCCTGGGAGGGTGTAAATTACAAAAAAATGAAAGTCGTTGCAAGGTCAATTAGGAGATGGC
 AAGGATATTGATATTCAAATAAGAAATTGATCCAAGGGTTGGTAAAGAGAACGCTGAAGTTCTATGAAAG
 ACCGTGTGAAGATTGCATGCAAAATCTAATAATATTAATAGAAATTAA

f488.aa

MPSSFPFLVNGSSGIAVGMATNMAPHNLREICDAIVYMLDNENASIFDLLKIVKGPDFPTFGEIVYNDNLIKAYK
 TGKGSVIRARYHIEERAEDRNAIIVTEIPYTVNKSALLMKVALLAKEEKLEGLLDIRDESREGIRIVLEVKG
 DPHVIMNLLYEYTEFKKHFSINNLALVNGIPKQLNLEELLFEFIEHRKNIIERRIEFDLRKAKEKAHVLEG
 NNIDEVIKIIKSSKLAKDARERLVSNFGLSEIQANSVDMRLQKLTALEIFKLEELNILLSLIKDYEDILLNP
 IINIIREETINLGLKFGDERRTKIYDEEVLKTSMSDLMQKENIVVMLTKKGFLKRLSQNEYKLQGTGGKGLSSFD
 LNDGDEIVIALCVNTHDYLFMISNEGKLYLINAYEIKDSSRASKGQNISELINLGQEEILTICKNSKDLTDDAYLL
 LTTASGKIAFESTDFKAVKSRGVIVIKLNDKDFVTSAEIVFKDEKVICLSSKKGSAFIFNSRDVRLTNRG
 QVGCGMQLKEGDLFVKVLSVKENP
 MKLKEGDLFVKVLSVKENPYLLIVSENGYKRLNMSKISELKRGATGTYTSYKSDKKAGSVVDAIAVSEDDEILL
 SKRSKALRTVAGKVSEQKDARGIQVFLDNDLSLVSVSKFIKZ

t488.aa

MATNMAPHLREICDAIVYMLDNENASIFDLLKIVKGPDFPTFGEIVYNDNLIKAYKTGKGSVIRARYHIEERA
 DRNAIIVTEIPYTVNKSALLMKVALLAKEEKLEGLLDIRDESREGIRIVLEVKGFDPHVIMNLLYEYTEFKKH
 SINNLALVNGIPKQLNLEELLFEFIEHRKNIIERRIEFDLRKAKEKAHVLEGNNIDEVIKIIKSSKLAKD
 AERLVSNFGLSEIQANSVDMRLQKLTALEIFKLEELNILLSLIKDYEDILLNP
 RRTKIIYDEEVLKTSMSDLMQKENIVVMLTKKGFLKRLSQNEYKLQGTGGKGLSSFD
 LNDGDEIVIALCVNTHDYL
 FMISNEGKLYLINAYEIKDSSRASKGQNISELINLGQEEILTICKNSKDLTDDAYLL
 LTTASGKIAFESTDFKAV
 KSRGVIVIKLNDKDFVTSAEIVFKDEKVICLSSKKGSAFIFNSRDVRLTNRG
 QVGCGMQLKEGDLFVKVLSVKENP
 YLLIVSENGYKRLNMSKISELKRGATGTYTSYKSDKKAGSVVDAIAVSEDDEILL
 SKRSKALRTVAGKVSEQGDARGIQVFLDNDLSLVSVSKFIKZ

f488.nt

ATGCCGTCATCATTCCATTCTTTGGAAATGGCTTAGTGGATTGCTGTGGAAATGGCTACTAATATGGCAC
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 ACTGGCAAGGGAAGTGTGTTATTAGGGCAAGATATCATATTGAGAAAGAGCAGAAGATAGAAATGCTATAATTG
 TTACAGAAATACCTTACGGTAAATACTGCACTCTTATGAAAGTGTGCGCTTGTGAAAGAGGATTAGGATAGTT
 AGAAGGACTTTAGATATAAGAGATGAATCTGATCGAGAAGGTATTAGGATAGTTCTGAAGTTAAAGAGGATT

TABLE 1. Nucleotide and Amino Acid Sequences

GATCCTCATGTTATTATGAATTGCTTATGAATATACTGAATTAAAAGCATTAGTATAAATAATTAGCCC
 TTGTTATGGTATTCCAAACAGTTAAATTAGAAGATTGTTATTGAATTATTGAGCATAGAAAAAATTATCG
 CGAAAGACGTATTGAATTGACTTGAGAAAGGAAAAGAGAAAGCACATGTTCTGAGGGATTAATATTGCTTA
 ATAATATAGATGAGTTATTAGATTATAATCATCTAAATTAGCAAAAGATGCAAGGGAGAGGCTTGC
 ATTTGGTCTTCAGAGATTAGGCCAATTAGTCTTGATATGAGTTACAAAACCTACAGCCCTGAGATT
 TAAGCTGAAGAGGAGCTTAATATACTGTTAAGCTTAATAAAAGATTAGAAGATATTCTCTGAACTCAGTAAGG
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 AGGTTCTTAAAGACTTCACAAAATGAGTATAAATTGCAAGGTACGGGAGGAAAGGACTAAGTCGTTGAT
 CTAATGATGGAGATGAGATTGTTATTGCTTGTGTCATAACTCATGATTATTATGATTCAATGAAG
 GAAAGCTTATTAAATCAATGCTTATGAAATAAAAGATTCTCAAGAGCTTCAAAAGGTCAAGAATATTAGGCT
 TATTAAATTAGGAGATCAAGAAGAAATATAACTATTAGAATAGTAAAGATTAACTGATGATGCTTATTG
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 TTATTAAACTGAATGATAAAGATTGTTACAAGTGCAGAGATTGTTTAAGGATGAAAAAGTAATTGCTTTC
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 ATGAAATTAAAAGAAGGTGATTGTTGTTAAAGTTTATCGTTAAAGAAAATCCTTATCTTTGATTGTTCTG
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 AGTAAACGTTCAAAAGCTTAAAGAACAGTAGCTGGAAAAGTATCTGAACAGCAAGATGCTAGAGGAATTCAAG
 TATTATTCTTGATAATGACAGCTGGTTCTGTTCAAAATTAAATAA

t488.nt

ATGGCTACTAATATGGCACCTCATATTAAAGAGAAATTGATGCCATTGTTACATGCTAGATAATGAGAATG
 CTTCTATATTGATTGCTTAAATAGTTAAAGGGCCTGATTCCAACTTTGGAGAGATTGTTATAATGATAAA
 TTTAATTAAAGCATAAAAATGGCAAGGGAAGTGTGTTATTAGGCAAGATATCATATTGAAGAAAGAGCAGAA
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 TAGCAAAGAAGAAAAGCTAGAAGGACTTTAGATATAAGAGATGAATCTGATCGAGAAGGTATTAGGATAGTTCT
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 AGTATAAATAATTAGCCTTGTAAAGGTTATTCCAAACAGTTAAATTAGAAGAATTGTTATTGAAATT
 AGCATAGAAAAAATATTATCGAAAGACGTATTGAAATTGACTTGAGAAAGGCAAAAGAGAAAGCACATGTTCTG
 GGGATTAAATATTGCTTAAATAATAGATGAGGTTATTAGATTATAATCTAAATTAGCAAAGATGCA
 AGGGAGAGGCTTGTGTTGAAATTGCTTCAAGAGATTAGGCCAATTAGTCTGATATGAGGTACAAAAC
 TTACAGCCCTGAGATTAAAGCTTGAAGAGGAGCTTAAATATACTGTTAAGCTTAAATAAGATTATGAGATAT
 TCTCTGAAATCCAGTAAGGATTATTAAATTATAAGAGAAACTATTAAATTAGGTTGAAATTGCGATGAA
 CGTCGAACTAAAATAATTATGATGAGGAGTTAAAAGCTAGTATGTCGGATTAAATGCAAAAGAAAATATTG
 TTGTTATGCTTACAAAGAAAGGTTCTTAAAGACTTCAAAATGAGTATAAAATTGCAAGGTACGGGAGGAAA
 AGGACTAAGTCGTTGATCTAAATGATGGAGATGAGATTGTTATTGCTTGTGTCATAACTCATGATTATT
 TTTATGATTCAAATGAAGGAAAGCTTTATTAAATCAATGCTTATGAAATAAAAGATTCTCAAGAGCTTCAAAAG
 GTCAGAATATTAGTGGACTTAAATTAGGAGATCAAGAAGAAATATTAACTATTAGAATAGTAAAGATTAAAC
 TGATGATGCTTATTGCTTACAACAGTGGAAAGATAGCTAGATTGCAATCTACAGATTAAAGCAGTA
 AAGTCACGGAGGTATTGTTAAACTGAAATGATAAGATTGTTACAAGTGCAGAGATTGTTTAAGGATG
 AAAAGTAATTGCTTCTAAAAGGGTAGTGCATTATTTAAATTCAAGGGATGTTAGGCTTACTAATAGAGG
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 TATCTTTGATTGTTCTGAAAATGGGTAAACATGCTAAATATCTGAGCTTAAAGAGGAG
 CCACTGGTTATACTAGTTATAAAAATCTGATAAAAAGCGGGTAGTGTGTTGATGCTATAGCAGTTCAGAGGA
 TGATGAAATCTGCTTGTAAAGTAAACGTTCAAAAGCTTAAAGAACAGTAGCTGGAAAAGTATCTGAACAGCAAA
 GATGCTAGAGGAATTCAAGTATTGTTCTGATAATGACAGCTGGTTCTGTTCAAAATTAAATAA

f494.aa

MFALIRKIFMIYFLCITLAGFAMIFIDSFKTEQPNVKENQSKINQHTIEPNLIMFTSSIGGFLGVYVGWIWFNYDK
 SNFYLNWGNLIIILYINIALIITVYSKSHS

t494.aa

TABLE 1. Nucleotide and Amino Acid Sequences

MIFIDSKFTEQPNVKENQSKINQHTIEPNLIMFTSSIGGFLGVYVGIFNYDKSNFYLNWGNLIIILYNTIALIIT
VYSKSHS

f494.nt

ATGTTGCATTAATTAGAAAAATTTATGATCTATTTTATGCATTACTCTGCAGGTTGCCATGATTTTA
TTGACAGCAAATTACCGAACAGCCTAATGTTAAAGAAAATCAAAGCAAATTAAATCAACATACAATTGAACCCAA
TTAATCATGTTACATCTATAGGAGGATTTAGGTGTTATGTTGGAATTGGATCTTAACATGACAAA
AGCAATTTCACCTAAATTGGGAAATTAAATAATATTAAACATAGCCCTAATTACTGTACTCAA
AATCACATAGTTAG

t494.nt

ATGATTTATTGACAGCAAATTACCGAACAGCCTAATGTTAAAGAAAATCAAAGCAAATTAAATCAACATACAA
TTGAACCCAATTAAATCATGTTACATCTATAGGAGGATTTAGGTGTTATGTTGGAATTGGATCTTAA
CTATGACAAAAGCAATTTCACCTAAATTGGGAAATTAAATAATATTAAACATAGCCCTAATTACT
GTATACTCAAATCACATAGTTAG

f516.aa

MKKTPNTCIFLTLLIISNLNALANEENNEKNDQPKQISNFFSPERGFIYSTGIGIGVGFFLNSNIKHLIFRPYY
TFSNNTFDLIVAMILTRESLNIPKKMQYFKSYIGGGINWHIANLIKTKYFSATIGIGGRFYLSTNFIEDIRFYE
KLPYVIEPYMFIEISSKKAIPLMGLDFKIDFLFLDTFNISFNFTIRYNFKDKNEMET

t516.aa

NEEGNTNEKNDQPKQISNFFSPERGFIYSTGIGIGVGFFLNSNIKHLIFRPYYTFSNNTFDLIVAMILTRESLNI
PKKMQYFKSYIGGGINWHIANLIKTKYFSATIGIGGRFYLSTNFIEDIRFYEKLPYVIEPYMFIEISSKKAIPLM
GLDFKIDFLFLDTFNISFNFTIRYNFKDKNEMET

f516.nt

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AAGGCAATACTAATGAAAAAAATGATCAACCCAAACAAATCTCAATTAGCCAGAAAGAGGGTTCATATA
TTCAACAGGAATTGGGATTGGAGTTGGATTCTAAATTCAAATATTAAACACCTTATCTTAGACCTTATTAT
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AAATGCAATACTTAAATCTTATATTGGAGGAGGAATAAACTGGCACATTGCAAACCTTAATTAAAAAAACAAAATA
TTTTCCGCCACCATTGGCATAGGTGGTCTTTTACCTATCTACAAACTTATAGAAAGACATTGATTTACGAA
AAATTGCTTATGTAATAGAGCCTTATATGTTATTGAAATTCTTCTAAAAGGCATTCTTAATGGGTTAG
ACTTAAATTGATTTTATTAGATACATTAAACATTCTTTAATTACTATTAGATATAATTAAAGGA
AAAAACGAGATGGAAACATGA

t516.nt

AATGAAGAAGGCAATACTAATGAAAAAAATGATCAACCCAAACAAATCTCAATTAGCCAGAAAGAGGGT
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f517.aa

TABLE 1. Nucleotide and Amino Acid Sequences

MIPVVASGGILIALSIAFVGIGPDGPNAEHFPYKQIADIGSIAFGMMLPVLAGFIAMAIADKPGLTPGLVGGVMS
 GNVKAGFLGAIIFAGFLAGYVARFLARRSVPWLRPVMPIFVIPLISTIIVGFFMLYFGVYIGKFMGVLESLKSLQ
 SNSETFGVLGKIFLGLVLGSMITVDMGGPFNKVAFLFGVGLIPQVPEIMGMVAAAIPVPPMAMGLATFLAPKLFEN
 EEKESGKIAFLISFIGISEGAIPFAASDPGRVIPSIVVGGAVSSIIAFLGVANHAPHGGPIVLVIDNKFGFIIA
 IAVGVAVATALVIFLKSLLKESE

t517.aa

DKPGLTPGLVGGVMSGNVKAGFLGAIIFAGFLAGYVARFLARRSVPWLRPVMPIFVIPLISTIIVGFFMLYFGVYI
 GKFMGVLESLKSLQNSNSETFGVLGKIFLGLVLGSMITVDMGGPFNKVAFLFGVGLIPQVPEIMGMVAAAIPVPPM
 AMGLATFLAPKLFENEKEKESGKIAFLISFIGISEGAIPFAASDPGRVIPSIVVGGAVSSIIAFLGVANHAPHGGP
 IVLPVIDNKFGFIIAIAAVGVAVATALVIFLKSLLKESE

f517.nt

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 GCTTGTGGTTATTGCAATGGCAATTGCTGATAAGCCTGGCTTACCCCGGTCTTGGGAGTAATGCT
 GGGAAATGTAAGCAGGTTCTTGGCGCAATTGCGGGCTTCTTGCAGGTTATGTTGCAAGGTTTAGCAA
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 CTTTTATGCTGTTGGGTATGAAATTGAGGCTTGGGCTTGGGCTTAAATCTTACAG
 AGTAATTGCAAACCTTGGCGTGGTAAAGCTTCTTGGGCTTGGGCTTAAATTGCTTAGGCTTAGTACTAGGTTCAATGATTACTGTTGATA
 TGGCGGACCTTTAATAAGTGGCATTCTTGGTGTAGGCTAATTCTCAAGTGCCAGAAATAATGGGAAAT
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t517.nt

GATAAGCCTGGCTTACCCCGGTCTTGGGAGTAATGCTGGAAATGAAAAGCAGGTTCTGGCGCAA
 TATTGCGGGCTTCTGCAAGGTTATGTCAGGTTAGCAAGAAGATCTGTTCTGAGTGGTTAGACCTGT
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 AAATTCTAGGCTTAGTACTAGGTTCAATGATTACTGTTGATATGGGCGGACCTTTAATAAGTGGCATTCT
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 ATAGTACTCTGTTATTGATAATAAATTGGGTTATTGCAATTGCTGTTGGAGTTGCGGTTGCAACAGCTT
 TGGTAATTGGAAATTCTTAAAGGAATCTGAATGA

f519.aa

MIKIFKKIYILTLVLMMAHLSFASDNYMVRCSKEEDSTTCIAKLKEIKEKKNYDLFSMGIGIGDPIANIMITIPYI
 NIDFGYGGFIGLKSNNFENYLNNGIDVFKKQIGQYMKIGGGIGIGADWSKTSLIPPNEEEETDYERIGAVIRIPF
 IMEYNFAKNLSIGFKIYPAVGPTILLTKPSILFEGIKFNFFGFGFIKFAFN

t519.aa

DNYMVRCSKEEDSTTCIAKLKEIKEKKNYDLFSMGIGIGDPIANIMITIPYINIDFGYGGFIGLKSNNFENYLNNG
 IDVFKKQIGQYMKIGGGIGIGADWSKTSLIPPNEEEETDYERIGAVIRIPFIMEYNFAKNLSIGFKIYPAVGPTI
 LLTKPSILFEGIKFNFFGFGFIKFAFN

TABLE 1. Nucleotide and Amino Acid Sequences

f519.nt

ATGATAAAAATTTAAAAAATACATTAAACATTAGTATTAGGTATGGCACACCTTCTTGACATCTGACA
 ATTATATGGTCAGATGCAGCAAGGAAGAAGATTCAACCACCTGTATCGCAAAGCTTAAAGAAATAAAAGAAAAGAA
 AAATTATGACTTATTTCAATGGCATTGGAATAGGAGATCCTATTGCAAATATTGATTACAATTCTTATATA
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 ACGTTATTTAAAAAGCAAATTGGACAATATGAAAATTGGCGCGCATTGGAATAGGTGCGGATTGGTCAA
 AACATCCCTTATACCCCTAATGAAAGAAGAAACTGATTATGAGAGAATAGGCCTGTTATAAGAATTCTT
 ATAATGGAATATAATTGCAAAAAATTATCCATAGGATTCAAATTATCCTGCAGTAGGGCAACAATTAC
 TAACAAAACCAAGCATTATTGAGGAATTCAATTGATTTGGATTGATTCATAAAATTGCAATTAA
 TTAA

t519.nt

GACAATTATATGGTCAGATGCAGCAAGGAAGAAGATTCAACCACCTGTATCGCAAAGCTTAAAGAAATAAAAGAAA
 AGAAAAATTATGACTTATTTCAATGGCATTGGAATAGGAGATCCTATTGCAAATATTGATTACAATTCTT
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 ATAGACGTTATTTAAAAAGCAAATTGGACAATATGAAAATTGGCGCGCATTGGAATAGGTGCGGATTGGT
 CAAAAACATCCCTTATACCCCTAATGAGAAGAAGAAACTGATTATGAGAGAATAGGCCTGTTATAAGAATTCC
 TTTTATAATGGAATATAATTGCAAAAAATTATCCATAGGATTCAAATTATCCTGCAGTAGGGCAACAATA
 TTACTAACAAAACCAAGCATTATTGAGGAATTCAATTGATTTGGATTGATTCATAAAATTGCA
 TTAATTAA

f520.aa

MRMLLATIILILTTGLAAQSKSKSMTEDDFDFDKLLAKEESVRLFGIGFGVGYPLANITISVPYVDIDLGYGGF
 VGLKPNNFLPYVVMGVDLLFKDEIHKNMISGGIGIGADWSKGSPEKSNEKLEEEENEAQVASLQNRIGVIRL
 PLVIEYSFLKNIVIGFKAVATIGTTMLLGSPMSFEGARFNFLGTGFIKIYI

t520.aa

QSKSKSMTEDDFDFDKLLAKEESVRLFGIGFGVGYPLANITISVPYVDIDLGYGGFVGLKPNNFLPYVVMGVDLL
 FKDEIHKNMISGGIGIGADWSKGSPEKSNEKLEEEENEAQVASLQNRIGVIRLPLVIEYSFLKNIVIGFKAV
 ATIGTTMLLGSPMSFEGARFNFLGTGFIKIYI

f520.nt

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 GTAGGGCTTAAACCAACAATTCTGCCCCATGTTGTAGGGTAGATCTCTATTAAAGATGAAATACACA
 AAAACACTATGATTTCTGGAGGCATTGGAATAGGTGCAGATTGTCAAAAGGAAGTCTGAAAAATCAAATGAAAA
 ACTTGAAGAAGAGGAAGAAATGAAGCACAACAGTAGCTTCTTCAAATAGAATAGGGTTGTGATAAGATTG
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 ATAG

t520.nt

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 TTTAAAGATGAAATACACAAACACTATGATTCTGGAGGCATTGGAATAGGTGCAGATTGGTCAAAGGAAGTC
 CTGAAAATCAAATGAAAACCTGAAGAAGAGGAAGAAATGAAGCACAACAGTAGCTTCTTCAAATAGAAT
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TABLE 1. Nucleotide and Amino Acid Sequences

f526.aa

MKKEFIMLLLLQTIMNLNSINTNTSTSIVKELQKNLYIFNSKEYQDKDTLNEFINSININDKEILQSLEKIKNE
 LFIISVFFNNKKGILIALNLGAEINFKYKISPISISIINNEFEITKILIDYGISLNQIDDTGYSPIFWAIYTNEK
 IFEFLKESGADLSFTLKNRKTpmQAAIETENIKLIKSLEKKKIYIDDNFKKKLKKLNKEIVRILVK

t526.aa

NSINTNTSTSIVKELQKNLYIFNSKEYQDKDTLNEFINSININDKEILQSLEKIKNLFIIISVFFNNKKGILIAL
 NLGAEINFKYKISPISISIINNEFEITKILIDYGISLNQIDDTGYSPIFWAIYTNEKIFEFLKESGADLSFTLKN
 RKTPMQAAIETENIKLIKSLEKKKIYIDDNFKKKLKKLNKEIVRILVK

f526.nt

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 TTTAAATGAATTATAAATTCAATAAAATATAATGACAAGAAATCTTACAAGTTAGAAAAAATCAAAATGAG
 CTTTTTATAATATCTGTTTTCAACAATAAAAGGGATTAAATTGCACTAAATCTTGGAGCAGAAATAAAACT
 TTAAATATAAATATCTCAATTCAATAAAACATGAATTGAAATCACAAAATATTGATAGATTA
 CGGAATAAGCCTTAATCAAATAGATGATACAGGTTATTCTCAATATTGGCAATATAACTAATAACGAAAAA
 ATATTGAAATTAAAAGAAAGCGGAGCTGATTAAAGTTCAACTTAAAATAGAAAAACACCAATGCAAGCCG
 CAATAGAAACAGAAATATAAAACTAATTAAATCTGGAAAAGAAAAAATTACATTGACGACAATTCAAAAA
 AAAACTAAAAAGCTAAAAACAAAGAAATAGTTCGAATTAGTAAATAG

t526.nt

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 AATATCAAAAAGATAAAGACACTTAAATGAATTATAAATTCAATAAAATATAATGACAAGAAATCTACAAAG
 TTTAGAAAAAATCAAAATGAGCTTTATAATATCTGTTTTCAACAATAAAAGGGATTAAATTGCACTA
 AATCTGGAGCAGAAATAACTTAAATATAAAATCTCAATTCAATAAAACATGAATTGAAA
 TCACAAAATATTGATAGATTACGGAATAAGCCTTAATCAAATAGATGATACAGGTTATTCTCAATATTGGC
 AATATATACTAATAACGAAAATATTGAATTAAAAGAAAGCGGAGCTGATTAAAGTTCAACTTAAAAT
 AGAAAAACACCAATGCAAGCCGAATAGAAACAGAAAATATAAAACTAATTAAATCTGGAAAAGAAAAAATT
 ACATTGACGACAATTCAAAAAAACTTAAAGCTAAAAACAAAGAAATAGTTCGAATTAGTAAATAG

f544.aa

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 LKEICVSILVGAILASVNFLRIVFFVAPHSDKLKIAFVVSCLMVLTVAKILGGLLPIVAKLLKLDPALMAGPL
 ITTIADAITLIAYFNIAKWVLVSYAV

t544.aa

STFTATIISNYQNLMLSLVVLANFIPLLMDTSGNAGSQASALIIRELALGTVVKDFFKVFLEICVSILVGAILA
 SVNFLRIVFFVAPHSDKLKIAFVVSCLMVLTVAKILGGLLPIVAKLLKLDPALMAGPLTTIADAITLIAYFN
 IAKWVLVSYAV

f544.nt

ATGACAAAAAATAGAATAATTGGCTTTAGTTCTTATGGTGTCTTCTACTTTACAGCTACAATTATTCAAATT
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 CTCTCAGGCATCTGCGCTAATAATTCTGTGAGCTTGCTCTGGTACTGTCAAGGTAAAAGATTTTAAAGTGT
 TTAAAGGAAATATGTGTTAGCATTCTAGTGGGAGCAATTCTGCTAGTGTAAATTAAAGAATTGTCTTTTG
 TAGCTCCACACCATTCTGATAAGCTGAAAATAGCTTTGTAGTTCATCTGCTTGATGGTAAGTTGACAGTAC
 AAAGATATTGGGAGGTCTTTACCCATTGTTGCTAAACTTTAAAGTTGATCCAGCACTTATGGCAGGCCCTTA

TABLE 1. Nucleotide and Amino Acid Sequences

GCTACTATTGGAACAACATATGCTACTTGGCAGCCCAATGTCATTGAAGGAGCTAGATTAAATTCTTAGGCACAG
GCTTTATAAAAATATATATAG

f523.aa

MNIKINFFFLPIGIFLGLFFPLGIYSSLSHAFIRLSYLSLIPFLIFIENIIENKNFKKLFKGKTIYYGILT
NLSGVAVSIIAATIYLPQRIPILEKTIQNTCFEKEALLETFFPKNIFKIFTSSNPNLLSIYMISIIIGTSFYAK
QKGRIARELMLSASNLFYHANGFIVNILNIGIIFITANYAANLKNFKDYPNYTNSITFFLAWTIIILFVILPTISY
RLTKSFKMIYKGIFVSFQNIIFSGLAKDSYSPYVILIEDIKNERINIKSIIINIPNLSVSKFGTIFVSVISFFI
ILKSYSSLPISIYEISYMSTLSFVFVFAFPHIPNSLIYIITMLCSTYTKGIELNVSNITPMLPILISLALLIDFAF
NIAIIHIINFKELKDQEKIN

t523.aa

IENIIENKNFKKLFKGKTIYYGILTNLSGVAVSIIAATIYLPQRIPILEKTIQNTCFEKEALLETFFPKNIFKIFT
SSNPNLLSIYMISIIIGTSFYAKQKGRIARELMLSASNLFYHANGFIVNILNIGIIFITANYAANLKNFKDYPNY
TNSITFFLAWTIIILFVILPTISYRLTKSFKMIYKGIFVSFQNIIFSGLAKDSYSPYVILIEDIKNERINIKSII
INIPNLSVSKFGTIFVSVISFFIILKSYSSLPISIYEISYMSTLSFVFVFAFPHIPNSLIYIITMLCSTYTKGIE
LNVSNITPMLPILISLALLIDFAFNIAIIHIINFKELKDQEKIN

f523.nt

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ATAGCTCCTTATCACATGCTTTATAAGATTATCATACTTATCTCTTATTCCCTTTAATATTTCAATTCCATT
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CAATACAAAATACATGTTTTGAAAAGAAGCTTACTAGAAACATTCTTCCAAAAAATATTTCAAAATATT
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CAAAAGATTCTTATTCCCTTATGTGATATTAAAGAGATATTAAACGAAAGATAAAATATAAAAAATCCAT
AATTATAAACATACCTTAATAAAATTGATCTAAATTGGCACTATTGTTCTAGTAATATCATTTCATA
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TCTTGCATTCCCTCATATAACCAAATAGTTAATTATATAATTACAATGCTTGCCTACATATAACAAAGGAAT
AGAGCTAAATGTTCAAACATAACACCAATGCTGCCGATATTAAATCTCTTGGCTTACTAATCGACTTGTCTT
AACATTGCAATCATTATATAAACTTCAAAGAATTAAAGATCAAGAAAAAATTAAATTAA

f523.nt

ATTGAAAATATTATTGAAAATAAAACCTTAAAAGCTTTGGTAAAACAATTATTATGGAATTAACTAAC
TATCTGGAGTTGCTGTATCAATAATAGCTGCAACAATATCTCCGCAAAGAATTCCAATACTAGAAAAACAAT
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ATTAATATAGGGATCATTTATAACAGCAAATTACGCTGCAAACCTTAAAGATTACCCAAATTAT
ACAAAACAGCATAACATTCTTTGGCATGGACAATTATAATTCTCGTAATATTGCAAACAATTAGTTATAGAT
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AGATTCTTATTCCCTTATGTGATATTAAAGAGATATTAAACGAAAGATAAAATATAAAAAATCCATAATT
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CTAAATGTTCAAACATAACACCAATGCTGCCGATATTAAATCTCTTGGCTTACTAATCGACTTGTCTTAAAC
TTGCAATCATTATATAAACTTCAAAGAATTAAAGATCAAGAAAAAATTAAATTAA

TABLE 1. Nucleotide and Amino Acid Sequences

ATCACTACAATTGCAGATGCTATTACTTAATAGCTTATTTAATATAGCAAAATGGGTTTAGCTATGCTG
TTTAA

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TCTACTTTACAGCTACAATTATTCAAAATTAAATGTTGCTTAGTGGTTTAGCTAATTTATTC
CCCTTTAATGGATACTCAGGCAATGCCGGCTCTCAGGCATCTGCGCTAATAATTCTGAGCTTGCTCTGGTAC
TGTCAAGGAAAAGATTTAAAGTGTAAAGGAAATATGTGTTAGCATTCTAGTGGGAGCAATTCTGCT
AGTGTAAATTAAAGAATTGCTTTGTAGCTCCACACCATTCTGATAAGCTGAAAATAGCTTTGTTAGTT
CATCTGCTTGATGGTAAGTTGACAGTAGCAAAGATATTGGGAGGTCTTACCCATTGTTGCTAAACTTTAA
GTTGGATCCAGCACTTATGCCAGGCCCTTAATCACTACAATTGCAGATGCTATTACTTAATAGCTTATTTAA
ATAGCAAAATGGGTTAGTTAGCTATGCTTTAA

f545.aa

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LKEICVSILVGAILASVNFLRIVFFVAPHHSIDLKIAFVVSSCLMVSLSVAKILGGLLPIVAKLLKLDPALMAGPL
ITTIADAITLIAYFNIAKWVLVSYAV

t545.aa

GSQASALIIRELALGTVVKVDFFKVLKEICVSILVGAILASVNFLRIVFFVAPHHSIDLKIAFVVSSCLMVSLSV
AKILGGLLPIVAKLLKLDPALMAGPLITTIADAITLIAYFNIAKWVLVSYAV

f545.nt

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TAGCTCCACACCATTCTGATAAGCTGAAAATAGCTTTGTTAGTTCATCTTGCTTGATGGTAAGTTGACAGTAGC
AAAGATATTGGGAGGTCTTACCCATTGTTGCTAAACTTTAAAGTTGGATCCAGCACTTATGGCAGGCCCTTA
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TTTAA

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TAATCACTACAATTGCAGATGCTATTACTTAATAGCTTATTTAAATAGCAAAATGGGTTTAGTTAGCTATGCTG
TGTAA

f577.aa

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YVSENLFYVISQINNVRFSFEKNKLNYNENQILKMLEYFSSFLNTKQMDLQKDFFNKGYGYLKLNKILLNKKSLIA
GLSDITFYNSLSEQEKSQIKFSYLIINDNNEIVISNPFIGILETSVLTKFINWILYKKTQKTLIGFNNQSQSNC
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t577.aa

NKNIVVLTDNKTIPFYINQFNIENKANFIKFRNNIDLQTIKEKENAQIIISKNIGNTNIANHFKSVKINYNPDYPI
LKHIFQFNYKIIPLGFDIPILYKNTHHIKKYINTKYLKEEYENFIKDGKFFISPYVSENLFYVISQINNVRFSF

TABLE 1. Nucleotide and Amino Acid Sequences

EKNKLNYNENQILKMLEYFSSFLNTKQMDLQKDFFNKYGYLKLNKILLNKSLIAGLSDITFYNSLSEQEKSQIK
FSYLIINDNNEIVISNPNFIGILETSVLTKFVINWILYKKTQKTLIGFNNQSNSNICFGFANGFTPYKELNLKIKHS
IDGISPFIIDETQINSHSYVLSKKTIEKENLLINEWFFSKANNLKKNN

f577.nt

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GTTAGAAATAATATTGATCTGAAACAATAGAAAAGAAAATGCACAAATAATTATTCTAAAAACATTGGTAAC
ACAAATAATTGCTAACCACTTAAATCTGAAAATCAATTATAATCCAGATTATCCTATCTTAAAGCATAATTCA
AGCAATTAACTACAAATTATCCATTGGGCTTGCACATTCTATTAACTATAAAATACACATCATATTAA
AAAATACATAAACACTAAATATCTAAAAGAAGAACAGAAAATTCTTAAAGATGAAATTTTATATCGCCT
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AAAAGATTCTTAATAAACCGCTACCTAAAGTTAAATAAAATATTGCTTAATAAAATCTCTTTAATAGCA
GGATTGAGCGATATAACCTCTACAATAGCTTAAGCGAACAGAGAAGTCACAAATAAAATTTCTATTAA
ACGATAACATGAAATTGTTATCTAACACCAATTATTGCTATTGCAATTAGAAACATCTGTTAACTAAAAAATT
TATCAACTGGATATTGTATAAAAAAACTCAAAACCTAATTGGATTAACTCAATCCAACTCAAATATGTT
TTGGATTGCAATGGTTTACCCCTACAAAGAATTAAATTAAAATAAACATTCAATTGATGGAATATCTC
CTTTATTATTGACGAAACTCAAATCAATAGCCATTCTATGTATTAGCAAAAAACATTGAAAGAAAACATT
ACTAATAAAATGAATGGTTTCTCAAAGCTAATAATCTAAAAAAATAAAATTAA

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AATAAGAACATCGTTGACTAACTGACAATAAAACAATACCATTATATAAATCAATTAACTGAAATAAG
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f584.aa

MIKTILLVLYPVVVFSQISANQYFEGIYAKYQNIEDMQATINFLKGLKQTGVLLYKFPDKFIINLDSNNQVFVS
DGEFLTVYVPSLGTFSNQQLLKSSGGGLMKVLNSEYSVSYTNSPNLEDLDSSEPGKYIKLTSRKLYKGAATINS
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PTSGGREIVIDLTAVKFNVGILDSDKFYDPPKSSNKVDNFLYDIKKN

f584.nt

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TABLE 1. Nucleotide and Amino Acid Sequences

AACAGGTGTTTGCTTATAAGTTCCAGACAAGTTATTATCAATTAGATTCAAATAATCAAGTTTGTAAGT
 GATGGTGAATTTTGACAGTTATGTTCCATCTCTGGGACTCTTTAATCAGCAATTATTAAGGGTAGTAGTG
 GGGGAGGCTTATGAAAGTTAAATAGTGAAGTATAGCTATCTTACCAATTCTCCAAATTAGAAGATCTCGA
 TTCATCTGAGCCTGGAAAATATTTAAACCTTTCTAGAAAGCTTACAAGGGGGCTGCTACTATTAAATTCT
 TTTATTATTGCTTTGCTCCGGATGGAATAATTAGAAGAATTACTGCTTTCTACTAGTGGTGGGCGCAAATAG
 TTATTGATTTGACTGCTGTGAAGTTAATGTTGGAATTCTGATAGCAAATTAAATGATCCTCCAAATCTTC
 AAATAAGGTAGATAATTAAATGATATTAAAAAAATTAA

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CAAATATCTGCAAATCAATATTGAAAGGAATTATGCTAAATATAGAGGACATGCAAGCAACAATTA
 ATTTTACTTAAAGGGGTTAAAGCAAACAGGTGTTGCTTATAAGTTCCAGACAAGTTATTATCAATTAGA
 TTCAAATAATCAAGTTTGTAAGTGAATTGAGCTTACAGTTATGTTCCATCTCTGGGACTCTTTAAT
 CAGCAATTATTAAGGGTAGTGTGGGGGAGGTCTTATGAAAGTTAAATAGTGAAGTATAGCGTATCTTACCA
 ATTCTCCAATTAGAAGATCTGATTCTGAGCCTGGAAAATATTAACCTTTCTAGAAAGCTTAA
 CAAGGGGGCTGCTACTATTAAATTCTTTATTGCTTTGCTCCGGATGGAATAATTAGAAGAATTACTGCTTT
 CCTACTAGTGGTGGCGCGAAATAGTTATTGACTGCTGTGAAGTTAATGTTGGAATTCTGATAGCAAATT
 TAAATATGATCCTCCAAATCTCAAATAAGGTAGATAATTAAATGATATTAAAAAAATTAA

f596.aa

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 YSKKYLFKNEHGVFFVKVNI PHGTSSIKYRLIVDGWTNDEYNKNVNVYNEFLIPFSKIEIAKEKSSYISLRN
 PIQSYDNNEIEIFYIGRPQIVTIAGSFNNFNPFLNRLEKEDNKGIVTIKLKNLPKDRIIYYYFIDSGNKVID
 KNNVNRINLYFVEGIDNKIDFEVSYFDHK

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DDYLIYDFDLSLNEFLEVSTRKDNLPMVDSNRILLFYPPKKEIRKIFAAFDQYSKKYL
 FKNEHGVFFVKVNI PHGTSSIKYRLIVDGWTNDEYNKNVNVYNEFLIPFSKIEIAKEKSSYISLRN
 PIQSYDNNEIEIFYIGRPQIVTIAGSFNNFNPFLNRLEKEDNKGIVTIKLKNLPKDRIIYYYFIDSGNKVID
 KNNVNRINLYFVEGIDNKIDFEVSYFDHK

f596.nt

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 GGATTAACTCCATTCTAAATTGAGATCGCTAAAGAGAAGTCCAGCTATATTCTTGAGAAATCCAATACAA
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GATGATTATTAATTATGACTTGTATTAAAGTTAAATGAATTCTAGAAGTTCAAACAAGAAAAGACAATCTG
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 AAATCCAATACAATCATGATAACAATGAAATTGAAATTTTTACATAGGTCGTCCTGGACAAATAGTTACAATA
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 TTAAGCTTAAATACCAAGGATAGAATTATTATTATTGATTCTGGTAACAAAGTAATAGATAAAA

TABLE 1. Nucleotide and Amino Acid Sequences

TAATGTTAATAGAATTAATTTATTTTGTGAGGGAATTGATAATAAAATAGATTTCGAAGTTCTATTTGAT
CATAAAGTAA

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MRQRVMIAAMALSCHPSLLIADEPPTALDVTIQEQLLILKNLSKKFNTSTIFITHDLAVVAEICDTVSVMYQGKIV
EEGTVEEIFNNPKHPTIGLLKSILTLEHDPNKKLYSTKENPMKITKTSTEEF

t598.aa

EPTTALDVTIQEQLLILKNLSKKFNTSTIFITHDLAVVAEICDTVSVMYQGKIVEEGTVEEIFNNPKHPTIGLL
KSILTLEHDPNKKLYSTKENPMKITKTSTEEF

f598.nt

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ATTTATAACTCATGATCTTGCGGTTGTTGCTGAAATTGTGATACAGTATCTGTAATGTATCAAGGAAAATTGTA
GAAGAAGGAACAGTAGAGGAAATATTAAACAATCCTAACGATCCTTACACCATTGGCTTTAAAATCAATTCTTA
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GGAGTTTAA

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ATACTCTACCATATTATAACTCATGATCTTGCGGTTGTTGCTGAAATTGTGATACAGTATCTGTAATGTATCA
AGGAAAATTGAGAAGAAGGAACAGTAGAGGAAATATTAAACAATCCTAACGATCCTTACACCATTGGCTTTA
AAATCAATTCTTACGCTAGAACACGATCCAATAAAAGCTTTATTCAACAAAAGAAAACCCTATGAAGATCACAA
AAACCAGCACCGAGGAGTTTAA

f600.aa

MAIMERSIIGLFIALAFVSWLTVARVVRGQVQSLSSSEFIQAAKTLGATNQRIILKHLIPNSIGMIVFTTIRVPS
FIMAEAFSLFLGLGISAPMTSWGEVQNGIATFVEYPWKVFIPAIVMТИFLFMNFLGDGLRDAFDPKDSI

t600.aa

RVVRGQVQSLSSSEFIQAAKTLGATNQRIILKHLIPNSIGMIVFTTIRVPSFIMAEAFSLFLGLGISAPMTSWGE
LVQNGIATFVEYPWKVFIPAIVMТИFLFMNFLGDGLRDAFDPKDSI

f600.nt

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AAGAATAATCTAAAACACTTGATCCCTAATAGCATTGGAATGATAGTTATTCACAACAATAAGGTTCCAAGC
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TGCAAAATGGAATTGCTACATTGTTGAATATCCATGGAAAGTTTATTCAGCTATAGTTATGACAATATTC
ATTATTTATGAACTTTAGGTGATGGCTAAGGGATGCTTGTGATCCAAAAGATAGCATCTAA

t600.nt

CGAGTTGTACGAGGCCAAGTACAATCACTATCAAGTTGGAATTTATACAAGCAGCCAAAACCCTGGTGCAACAA
ATCAAAGAATAATCTAAAACACTTGATCCCTAATAGCATTGGAATGATAGTTATTCACAACAATAAGGTTCC
AAGCTTTATTATGGCTGAAGCATTTCATCCTTTAGGACTTGGAAATTTCAGCTCCAATGACAAGCTGGGAGA

TABLE 1. Nucleotide and Amino Acid Sequences

TTAGTGCAAAATGGAATTGCTACATTGTTGAATATCCATGGAAAGTTTTATTCCAGCTATAGTTATGACAATAT
TTCTATTATTATGAACCTTTAGGTGATGGCTAAGGGATGCTTGTCAAAAGATAGCATCTAA

f603.aa

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PSLKKKDLTVSQYIKLGFPKSLTLGVISLIISLSIGIPIGILAAIYKNTYVDIITSIAILGISIPLFVIGPILQY
FFAIKWGLLYTSGWITERGGFSNLILPIITLSMPNVAIFARIIRGSMLEIIQSDFIRTARAKGLSFKKIVIKHMLR
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RV

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)
SPFDSEKPIDPQVKARLMEKYHLDKPFYIQAFYYITNALRGDLGSPSLKKKDLTVSQYIKLGFPKSLTLGVISLIIS
LSIGIPIGILAAIYKNTYVDIITSIAILGISIPLFVIGPILQYFFAIKWGLLYTSGWITERGGFSNLILPIITLS
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MGMFITESALNRDYPVLMGGLVYSIILLISILISDIYKILDPRV

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AAAATATCACCTTGACAAGCCTTTATATTCAAGCTTTTATTACATTACAAACGCTCTCAGGGGAGATCTGGGA
CCTCTTGAAAAGAAAGACCTTACAGTTAGTCATAACATAAAATTAGGATTCCAAAATCACTACACTAGGAG
TAATATCCCTTATTATTCACATCAATAGGAATACCAATAGGTATATTAGCTGCCATTATAAAACTTATG
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TTTTTGCAATTAAATGGGTTGCTTATACCTCTGGATGGATTACAGAAAGAGGAGGATTCAAATTAAATTC
TACCCATAATAACTCTAGCATGCCAACGCTAGCTATTTCGCAAGAATAATCAGAGGATCAATGCTAGAAATAAT
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GGAGCAATGTTGCCTGTAGTAAGCTATATAGGTCCAGCATTGCTGCTATAATATCTGGAAGCGTGGTTATTGAAA
AAATATTAGAATTGCTGGAATGGGATGTTATAACAGAACCGCACTAAACAGAGATTACCCAGTATTATGGG
CGGATTGTTAGTATATTCAATAACTGCTTATTCTATATTAAATCAGATATTATATAAAATATTAGATCCA
AGAGTATAA

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AGTCCATTGATTCTGAAAACCTATTGATCCTCAAGTAAAGCAAGATTGATGGAAAATATCACCTTGACAAGC
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ATGGGAATGTTATAACAGAACCGCACTAAACAGAGATTACCCAGTATTAAATGGCCGGATTGTTAGTATATTCAA
TAATACTGCTTATTCTATATTAAATCAGATATTATATAAAATATTAGATCCAAGAGTATAA

f607.aa

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KSWNISEDIIYTFNLREDIVWSDGVITAEEIKKSYLRILNKTKAAMYANLIKSTIKNAQEYFDETVPESELGIK
AIDSKTLEITLTSPKPYFPDMLTHSAYIPVPMHIVEKYGENWNPENIVVSGAYKLKERSINDKIVIEKNEKYNA
KNVEIDEVIFYPTEGSVAYNMYINGELDFLQGAEKNNLLEEIKIRDDYSGLNGMAYIAFNTTIKPLDNLKVRQAI
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TABLE 1. Nucleotide and Amino Acid Sequences

FLQEQQKKILNINLEIENEETFLGSRTGNYQMSSVGWIGDYFDPLTFLDSLFTTENHFLGAYKYSNKEYDALI
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CISNAKKEKIVFRVSNLSEPSSLDPQLSTDLYGSNIITNLFLGLAVKDSQTGKYKPGGLAKSWNISEDGIIYTFNLR
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FPDMTHSAYIPVPMHIVEKYGENWTNPENIVVSGAYKLKERSINDKIVIEKNEKYYNAKNEIDEVIFYPTEGSV
AYNMYINGELDFLQGAEKNLLEEIKIRDDYSGLNGMAYIAFNTTIKPLDNLKVRQAISLAIDRETLTKVVLKGS
SDPTRNLTPKFDDYSYGKNNLILFDPEAKLLAEAGYPDGKGFPTLKYSKISEGRPTTAEFLQEQFKKILNINLEIE
NEEWTFLGSRTGNYQMSSVGWIGDYFDPLTFLDSLFTTENHFLGAYKYSNKEYDALIKKSNFELDPIKRQDILR
QAEIIIAEKDFPMAPLYIPKSHYLFRNDKWTGWVNPNAESYLYEDIKK

t607.nt

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TTTAATAAAATCTACAATAAAAATGCACAGAACATTTCGATGAGACAGTGCCTGAATCTGAGCTTGGCATAAAG
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ATTATTGATCCTTAACATTCTTAGACAGCTTATTACAACAGAAAATCATTAACTTAGGAGCGTACAAATATTG

TABLE 1. Nucleotide and Amino Acid Sequences

AAACAAAGCTATGCTTAATAAAAAAATCTAATTTGACTTCAATAAAAGACAAGACATTAAAGA
 CAGCTGAAAGAGTAAATAGCAGAAAAAGACTTCCTATGGCACCTTATATATACCCAAATCTCATTATCTTCA
 GAAATGATAATGGACACGGTGGTACCAAATATCGCAGAAAGCTATTTATATGAAGATATTAAAACAAAAATA
 A

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MKYLFLFLFISFTLFGFEDSSLKIGIDDVYVEAHEEGFHLFIRKKPAIKSVILTESFEIPDKKKDVATYSFRTLSY
 NKVAGDEIPILNGRVIKNKELLSLTSSTPVNKKFGEAFHILIPKKLKYGFPNFSTRSGDIDLEVLKSKEFWFS
 IFSFEKKYNDYLGRYQDNAYELLFKDDQNQGKIEFNELKDTFTKSDEVVIANNGIDIVDKINKILKNSEDSVYDL
 DLVLTVDVTDMSKSNIEILKEHLSIIIEPQLQKFKSYRIGLVFYKDYLEDFLTKAFDFNTIPYLNNILKYVNNGGG
 GDYPEAVFEGIDAATQFDWRAERRFIIVIGDAPPHEYPRGSIVYKDVINSAKEKDITIYGIIFQ

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FEDSSLKIGIDDVYVEAHEEGFHLFIRKKPAIKSVILTESFEIPDKKKDVATYSFRTLSYNKVNDEIRILNGRVI
 KVKELLSLTSSTPVNKKFGEAFHILIPKKLKYGFPNFSTRSGDIDLEVLKSKEFWFSIRSFEKKYNDYLGRYQ
 DNAVYELLFKDDQNQGKIEFNELKDTFTKSDEVVIANNGIDIVDKINKILKNSEDSVYDLVLVVDVTDSMKSNI
 EILKEHLSIIIEPQLQKFKSYRIGLVFYKDYLEDFLTKAFDFNTIPYLNNILKYVNNGGGDYPEAVFEGIDAAVT
 QFDWPAERFPIIVEGAPPHEYPRGSIVYKDVINSAKEKDITIYGIIFQ

f611.nt

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 ATAAAGGTTATGGAGATGAAATTCGGATTAAATGGAAGAGTTATAAGAATAAAGAACCTTTATCATGACAT
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 TCCAAAGGAAAGATATTACAATTATGGAATAATATTCACTGAA

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TTGAAAGATAGTTCTTGAAAATAGGTATTGATGATGTTATGTTGAGGCTCATGAAGAGGGATTCATCTTTTA
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 TCTTGTGTTATAGATGTTATCAATTCTGAAAGAAAAGATATTACAATTATGGAATAATATTCACTGAA
 CTATTGTTATAGATGTTATCAATTCTGAAAGAAAAGATATTACAATTATGGAATAATATTCACTGAA

TABLE 1. Nucleotide and Amino Acid Sequences

f617.aa

MIFFRNSFMALIFSFSILSISYFFGDFQFSYIKMISWRFILFLIMATGIATCAKSNSLNLNEGQIYFGAFLVYI
 FSSFFGLTYFNFVFLILLSSFFVGLGLIPFFITFFFGLNKALTGLLISYGNQRLVDGFLINMLKTGSFSNQTKRI
 NSLFALDSSLIYLFLLGVSWLFYVFIHKKTIYGLQLEILSNKKIDIFFNINEFKYKFFAVFGSAFLNGLAGSMF
 VVFFRPyVLGLTSGLWSSLIVAVISGFNYVYVLFSSLFSILIEFNNFLNINYDFKYEFIGLCQSLIAIFISLFL
 IKARKK

t617.aa

AKNSLNLNEGQIYFGAFLVYIFSSFFGLTYFNFVFLILLSSFFVGLGLIPFFITFFFGLNKALTGLLISYGNQ
 RLVDGFLINMLKTGSFSNQTKRINSLFALDSSLIYLFLLGVSWLFYVFIHKKTIYGLQLEILSNKKIDIFFNIN
 EFKYKFFAVFGSAFLNGLAGSMFVFFRPYVLGLTSGLWSSLIVAVISGFNYVYVLFSSLFSILIEFNNFLNI
 NYDFKYEFIGLCQSLIAIFISLFLIKARKK

f617.nt

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 TACTTGTGCCAAGAGTAATTCAATTAAATCTTGGGAATGAAGGTAGATTATTTGGGGCATTAGTTATATA
 TTTCAAGTTTTTGGATTAACCTATTAAATTGGTATTTTGATACTTTAAGTTCTTGTAGGACTTT
 TGGGGCTATCCCCTTTTATTACTTTCTCGATTAAATAAGCCTTAACAGGTCTTTAATATCTTATGG
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 TTATTACAAAAAAACTATTATGGCTTCAGCTTGAATATTAAAGCAATAAAAAAGATAGACATTTTCAA
 TATAAAATGAATTAAATATAAGTTTCGCTGTATTGGCAGTGCCTTTAAATGGCTTGCAAGGTTCTATGTT
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 TTTCAGGATTTAATTATGTTATGTTATTAGCTTATTGTTCAATATTAAATGAATTAAATAATTCT
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 AATTATGACTTAAGTATGAATTATTGGGCTTGTCAATCAATTGCTATTATCTCTTATTGTTGATTAAAG
 CTAGGAAAAAGTAG

f631.aa

MVVEINSLRTCYLLVLLLVAYGLVVFYTSFFLSLELTGNPNFLFFTRLNYLFLSFMVFLVFERISLNFLKKSIF
 PVLIITLFLIMATFLSPSISGAKRWIFFQGVSIQPSEIFKISFTIYLSAYLSKFDPRKNNGISYWIKPMLIFW
 VLIILQNDYSTAIYFAILFFIVLFVSNMAFSYVFAIVVTFLPVSAILMLEPYRVSRIFAFLNPYDDPSGKGYQII
 ASLNALKSGGILGKGLGMGEVKLGKLPEANSDFIFSVLGEELGFLGVLFIAISLFFLFYFGYFIAHSNSRFKFFI
 AFISSLAIFLQSMNNILIAIGLLPPTGINLPFESSGGSSIIVTMALSGLISNVSKNLSNN

t631.aa

TABLE 1. Nucleotide and Amino Acid Sequences

RISLNFLKKSIFPVLIIITLFLIMATFLSPSISGAKRWIFFQGVSIQPSEIFKISFTIYLSAYLSKFDPRKNNGISY
WIKPMLIFAIFWVLILQNDYSTAIYFAILFFIVLFSNMAFSYVFAIVTFLPVSIFLMLPEYRVSRIFAFLNP
YDDPSGKGYQIIASLNALKSGGILGKGLGMGEVKLGKLPPEANSDFIFSVLGEELGFLGVLAISLFFLFFYFGYFI
AIHSNSRFKFFIAFISSLAIFLQSMMNILIAIGLLPPTGINLPFFSSGGSSIIVTMALSGLISNVSKNLSNN

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ATGGTTGAGAGATAAATTCACTTAGGACATGTTTGTGCTATTGGTAGCCTATGGCCTTGTAG
TTTTTATACTTCTTCTTTCTAAGCTTAGAATTGACAGGTAATCCAAATTTCACAAGACTTAA
TTATCTTTTTAAGTTTATGGTTTCTGTTGAAAGGATTCTTAAATTTCAGGAAATCAATATTT
CCTGTATTGATTATAACTCTTTAATTATGGCAACTTTTATCTCAAGTATTCTGGAGCAAAGAGATGGA
TATTCTTCAGGTGTTAGCATTCAACCTCTGAGATTAAAATATCTTTACTATTTATCTTCAGCTTATT
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ACCTTATAGGGTTCTAGAATTTCGCCTTCTCAATCCTACGATGATCCTCTGGCAAAGGTTACAGATAATA
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TACCAAGGCCAATTGGATTATTTCAGTTCTGGAGAAGAATTAGGATTAGTTAGGGTTTGTGCTAT
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CAGGGATAAATTACCATTTTCATCTGGGGATCTCTATTATGTTACCATGGCATTGCTGGCCTTATTTC
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TGAATATTCAATTGCAATCGGTCTTGCCTCCTACAGGGATAAATTACCATTTTCATCTGGGGATCTTC
TATTATTGTTACCATGGCATTGCTGGCCTTATTCAATGTTCAAAAATTAAAGTAATAATTGA

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MKVNNFLSFFFRAFFLLFLIVILFFFVLFIDFIGMYNTKRYFPEFVRTKLLGETSLVFDHNSNIILDEARLVKER
EAIDIKNQQIEKLKEDLKLKEDSLNKLFEFELKQKQKDLQKQIIDDIINKYDEEANILQTAVALMNMPPEAVK
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PYWLSLMDSKKAAILIRKMSVSSLE

f647.nt

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GTTAGGAGAAACTTCTCTGGCTTTGATCATAATTCTAATATAATTCTGATGAAGCTAGACTTGTGAAGGAAAGA
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TABLE 1. Nucleotide and Amino Acid Sequences

AGCTTGAATTGAGCTTAAGCAAAAGCAGAAAGATTAGATTTAAAACAAAAATAATAGATGACATTATAAAATAA
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CGGCTTGAAGATTAAATCCCGAGCTTGCATATCTATATGCCGAAAATTGAAGAGCTTCCAAAAAGAAGGTC
GTTTATCAATTGTTCTTATTGGTTATCTTATGGATTCTAAAGCTGCTATATTGATTAGAAAAATGTCTGT
TAGTTCATTGGAGTAG

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CTCTGGTCTTGATCATATTCTAATATAATTCTGATGAAGCTAGACTTGTGAAGGAAAGAGCTATTGATAT
TAAGAACATCAGCAGATTGAAAAGCTTAAAGAAGATCTAAAGTTAAAGAACAGCTTAAATAAGCTGAATTGAG
CTTAAGCAAAAGCAGAAAGATTAGATTAAACAAAAATAATAGATGACATTATAAAATAATGATGAGG
AAGCAAATATTTGCAAACAGCTGTATATTAAATGAATATGCCACCAGAAGATGCTGTTAGCGGCTTGAAGATT
AAATCCCGAGCTTGCATATCTTATATGCCGAAAATTGAACAGCTTCCAAAAAGAAGGTCGTTATCAATTGTT
CCTTATTGGTTATCTTATGGATTCTAAAGCTGCTATATTGATTAGAAAAATGTCTGTTAGTCATTGGAGT
AG

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MLTYGDMVTLVFFVTMFSLNDIIFQENVIRIMSASFTGAGFFKGGKTLDFSKLSYLSNSFMSLPSTVRNKQASQ
TAKNKS MIEFIEKIQSKNIVRQEERGIVISLAADAFFDSASADVKLEENRDSIQKIASFIGFLSPRGYNFKIEGH
TDNIDTDVNGPWKSNWELSAARSVNMLEHILNYLDQSDVKRIENNFEVSGFGGSRPIATDDTPEGRAYNRRIDILI
TTDASLSFPKEIKQ

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NDIIFQENVIRIMSASFTGAGFFKGGKTLDFSKLSYLSNSFMSLPSTVRNKQASQTAKNKS MIEFIEKIQSKNIVV
RQEERGIVISLAADAFFDSASADVKLEENRDSIQKIASFIGFLSPRGYNFKIEGHTDNIDTDVNGPWKSNWELSA
RSVNMLEHILNYLDQSDVKRIENNFEVSGFGGSRPIATDDTPEGRAYNRRIDILI TTDASLSFPKEIKQ

f653.nt

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ACAAGCATCTCAGACTGCTAAAATAATCCATGATTGAAATTATTGAGAAGATTCACTAAATATTGAGTT
AGGCAAGAAGAAAGAGGTATTGTAATATCTCTGCAGCAGATGCATTGGTATTGCTAGTCAGATGTTAAGC
TTGAAGAGAAATAGAGATTCTATTCAAAAATAGCATCTTATTGGCTTTAAGTCCTAGAGGCTATAATTAA
AATAGAAGGGCATACAGATAATATTGATACTGATGTAATGGACCTTGAAAAGCAATTGGAACTTCGGCTGCT
AGATCTGTTAATATGCTGGAACATATTGAACTATTGATCAATCTGATGTTAAAGAATTGAAAATAATTG
AAGTATCTGGTTTGGTGGAAAGTAGGCCTATTGCAACAGACGATACCCCTGAGGGTAGGGCTATAATAGAAGAATT
TGATATATTAAATTACTACAGATGCATTTAAGTTCCCTAAGGAAATTAGCAGTAA

f664.aa

TABLE 1. Nucleotide and Amino Acid Sequences

MRMSVYTMGFAYIRSIMGYVVLFFFASLAVNFFVNIIQVGFFITFKSLEPRWDKISFNFSRWAKNSFFSAGAFFNL
FKSLLKVVICLIIYFIENNIGKISKLSEYTLQSGISIVLVIAYKICFFSVMFLAIVGVFDYLQFQRQYIESLKM
TKEEVKQERKEMEGDPLLRSRIKERMVILSTNLRAIPQADVITNPEHFAVAIKWDSETMLAPKVLAKGQDEIA
LTIKKIARENNEVPLMENKLLARALYANYKVNEEIPREYWEIVSKILVRVYSITKKFN

t664.aa

FVNIIQVGFFITFKSLEPRWDKISFNFSRWAKNSFFSAGAFFNLFKSLLKVVICLIIYFIENNIGKISKLSEYTL
QSGISIVLVIAYKICFFSVMFLAIVGVFDYLQFQRQYIESLKM TKEEVKQERKEMEGDPLLRSRIKERMVILST
NLRAIPQADVITNPEHFAVAIKWDSETMLAPKVLAKGQDEIA LTIKKIARENNEVPLMENKLLARALYANYKVNE
EIPREYWEIVSKILVRVYSITKKFN

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GTTGGATAAAATTACTTTAATTTTCCAGATGGCAAAAAAATTCTTTCTCAGCAGGGGCTTTTCAATTG
TTAAAAAGTTGTTAAAAGTTGTTAATATGCTGATATTATTATTTATATAGAAAACAATATAGGAAAATT
CTAAGCTTCGGAGTACACTCAATCTGAATTCTCTATTGTTAGTGCCTATAAGATATGTTTTTC
AGTAATGTTTGGCAATTGTAAGGGGTGTTGATTATTGTTCAAAAGATCTCAGTACATTGAGAGTTGAAAATG
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TGAGGGTTATTTAAGTACCAATTAAAGAGTAGCTATTCTCAAGCAGATGTAGTAATTACAAATCCAGAACATT
TGCAGTTGCTATTAAATGGGATAGCAGAACATGTTAGCTCCAAGGTGCTGCAAAGGTCAAGATGAAATAGCT
CTCACAAATTAAAAATTGCAAGAGAAAATAATGTTCTTAAAGCTCCTTGCAAGAGCTCTTATG
CTAATGTTAAGGTTAATGAAGAGATTCCAAGAGAATATTGGGAGATTGTTCAAAATTCTGTGAGAGTATATT
TATTACTAAAAAGTTAATTAG

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TGTATAATATGCTGATATTATTATAGAAAACAATATAGGAAAATTCTAAGCTTCGGAGTATACA
CTCAATCTGAATTCTATTGTTAGTGCCTATAAGATATGTTTTTCAGTAATGTTTGCAATTG
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AAGAGAAAATAATGTTCTTAAAGGAAAATAAGCTCCTTGCAAGAGCTCTTATGCTAATGTTAAGGTTAATGAA
GAGATTCCAAGAGAATATTGGGAGATTGTTCAAAATTCTGTGAGAGTATATTACTAAAAAGTTAATT
AG

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SFLSLIPIVFGAILLKHKEFYDIFMVLNFFEINLGALVAFVVGIFSINFFKMLNNKKLYYFSIYLFALSIIVCYF
VRI

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ITGILILMLEFNFLKVDFKGNILLAGIFMGLMQGLGALPGISRSGITIFSASVIGFNRKSAFEIISFLSLIPIVFGA
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VRI

f680.nt

TABLE 1. Nucleotide and Amino Acid Sequences

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 AGGAATCTCGTTCAGGAATTACGATCTTCGGCATCGGTTATGGATTTAATAGAAAAAGTCATTTGAAATT
 TCATTAAATTCTTAATTCCAATAGTTGGAGCGATTAAACATAAAGAATTATGATATTAAATTG
 TTTAAATTGAAATAACAAACTTAGGAGCATTAGTTGCTTGTGTTGGTATTTCTCAATAAAATTCTTTT
 TAAATGCTTAATAACAACAAACTGTATTATTTCTATATATTATTTGACTTCAATTATAAGTTGTTATT
 GTAGAATATGA

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 ATCGGTTATGGATTTAATAGAAAAAGTCATTTGAAATTTCATTAAATTCCAATAGTTGGAGCG
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 TTGCTTGTGTTGGTATTTCTCAATAAAATTCTTTAAATGCTTAATAACAAAAACTGTATTATTTTC
 TATATATTATTTGCACTTCAATTATAGTTGTTATTGTTAGAATATGA

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 GLKIEINFDEKDYFESTSVKTLNLMQEMGGISIFKIEIEGRPGEFKNAKAMQILDLITDKLDAFSAKTQSS
 INGILKFTNFKIKKESPLEYKLPENKIIILNKLINLIDKSDWTKDNKRMYINDDWSLISIIVRIEDNSTEGIKKFEK
 YAINTINEYMKNNKYHFSGVYDKVLIAKTMVKEQVINIITLGSITLLMFFFKSIKTGIIIAIPVAWSVFLNFAV
 MRLFGITLNPATATIASVSMGVGVVDYIHFNTFILQYQKNQIYKTALLESIPNVFNGIFANSISVGIGFLTLFS
 SYKIIISTLGAIIAFTMLTTSLASLTLPLLIYLFKPRVKLASNNNFKKLKQZ

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YKTMGIFMSIGVIISMIISLTVPGLIITLIPFAKKSFKEKEKENKLNKISFLERLAKLNTQITKSILKRKYTSSIM
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 AFSAKTQSSSINGILKFTNFKIKKESPLEYKLPENKIIILNKLINLIDKSDWTKDNKRMYINDDWSLISIIVRIEDN
 STEGIKKFEKYAINTINEYMKNNKYHFSGVYDKVLIAKTMVKEQVINIITLGSITLLMFFFKSIKTGIIIAIPV
 AWSVFLNFAVMRLFGITLNPATATIASVSMGVGVVDYIHFNTFILQYQKNQIYKTALLESIPNVFNGIFANSISV
 GIGFLTLTFSYKIIISTLGAIIAFTMLTTSLASLTLPLLIYLFKPRVKLASNNNFKKLKQZ

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 TATGCTATTAAACACAATTAGAAATATGAAAAATAATAATATCATTCTCAGGTGTTATGATAAGGTATTAA
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 ATTCAATTCTATTTCATACATTATTACAAATACCAAAAAATCAAATCTACAAAACACTGCACTTCTGAAATC
 AATACCCAAATGTATTAAATGGAATATTGCAAATTCTATTCTGTTGAATAGGATTAACTCTAACATTTCG

TABLE 1. Nucleotide and Amino Acid Sequences

TCTTATAAAAATAATCAACTCTGGAGCAATAATTGCTTTACAATGCTAACGACATCTCTGCATCACTAACTCTCTTCCATTATTAATTTATTAAACCTAGAGTAAAGCTAGCCTCAAACAACAATTAAAATTAACATAAA

t688.nt

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f704.aa

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t704.aa

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f704.nt

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t704.nt

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TABLE 1. Nucleotide and Amino Acid Sequences

AAAACTTTACATTACATTGTAGCACAAATATTAGGAGCTTTACAGGTGCATTAATGACACTTGTGCTATTTAT
 CCTAAATGGATAGAAATGGATCCTGGCTTAGAAAATACTCAAGGAATAATGGCAACTTCCCTGCTTCCCTGGAT
 TTTGCCTGGATTTATTGATCAAATTGGAACTTTGCTAATGTTTTAATTCTGTTGGAGATTTAC
 AAAAAACACAGCGACAATCCATTATTCTTTATTGCTAGGAGCAGTGGTTATCAATAGGGATAAGTTCGGA
 GGAATGAACGGTTATGCTATTAATCCTGCAAGGGATCTGGGACCAAGAATTTACTCTATTGCTGGATTTAAA
 ATCACGGATTAACAATCTAAGTATAGTTATTGCTACCAATAATTGGCCAATAATTGGAGCAATTGGAGGCTAC
 AATTACGAATTACACTAAAAATAACAAAG
 ACTAA

f707.aa

MRRLFLLYILCSFVFLNLFAQGSSSYIDKQKELAIFYYEVGQRYINVGKIKKGKLFQAKALKIYPDLKKGFDIKLA
 VKELDARIKDDNPKVVMLEDIKLEIIPGIVHEKIEINDFTNAPKIEYIAQRERSKNQDKIIKFQFGKFARALISRN
 FDLFDSVIADKVNVMQFESKNDFISTLSSASSKADADELEYLSVDDYYDLKSLKISKNSDTSFAVNVNNAKKNDVT
 KNFPFWKERQTLIFTTEDNNWFLOSSINZ

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MRRLFLLYILCSFVFLNLFAQGSSSYIDKQKELAIFYYEVGQRYINVGKIKKGKLFQAKALKIYPDLKKGFDIKLA
 VKELDARIKDDNPKVVMLEDIKLEIIPGIVHEKIEINDFTNAPKIEYIAQRERSKNQDKIIKFQFGKFARALISRN
 FDLFDSVIADKVNVMQFESKNDFISTLSSASSKADADELEYLSVDDYYDLKSLKISKNSDTSFAVNVNNAKKNDVT
 KNFPFWKERQTLIFTTEDNNWFLOSSINZ

f707.nt

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 CCATAAAATTGA

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f709.aa

MLIFGFIGLFFLNIFSLHAQGIVTNKDAQEEFKWALNSYNNGIYDDALLSFKKILSFDPNNLDYHFWTNVYYRLG
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 YAANFVGNEILYFDVNNNVNALVKDGFSYLKSPYDIEANNLYVTLYSDEIGVYDKVLGVKRKSIGNKGTDGE
 LLAPQYMAIDKRNYIYVSEWGNKRVSKFLEGDFILHFGSRSGYKGLLGPVTYLNENIYVADSLRNTIEVFDT

TABLE 1. Nucleotide and Amino Acid Sequences

SGNHLYSVFTSIEGIEGLSSDFVGNNVIVSSKGVDVYKYSIAKKTITKILKADKMNSKISSSSILDANNQMIVSDFNN
AKVSVYKSDASLYDLSNVDVRIIIRLGGPKIYVELNVSSKGLPVVGLKSENFISISNENYYIVNPKVAYNVNASKD
INIAVVFDFDKSSYMKYDTDQIVGLNAMLMELSKNKNFSFINATSVPIIDNIESLTSIRNTSSLGPYSTDAVKTDV
LKLAGSGLMSKSSRRAVVYFSGGILNRKAFEKYSLDTIVSYYKNNDIRFYLILFGNDPINSKLQYLVNETGGAVIP
FSSYEGVSKVYDLILEQKTTGTYLLEYYYPGPQEPNKYFNLSSVEANINQQTGRGEFAYFIN

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QGIVTNKDAQEFKWALNSYNNGIYDDALLSFKKILSFDPNNLVDYHFWTGNVYRLGYVEEALMEWRNLKDQGYKV
PYLRHLISTIEQRGIFSNEYLFKKLVVASLDNSIYKRPHGYQITSLRADKYGGYYAANFVGNEILYFDVNNNV
NALVKDGFSYLKSPYDVEANNLVTLYSSDEIGVYDKVLGVKRKSIGNKGTDGEALLAPQYMAIDKRNYIYVSE
WGNKRVSKFGLEGDFILHFGSRTSGYKGLLPGTVTLNENIYVADSLRNTIEVFDTSGNHLYSVFTSIEGIEGLS
SDFVGNNVIVSSKDGVYKSIAKKTITKILKADKMNSKISSILDANNQMIVSDFNNAKVSVYKSDASLYDSLNU
VRRIIIRLGGPKIYVELNVSSKSGLPVVLKSENFSISNENYYIVNPKVAYNVNASKDINIIVVFDKSSYMKKYD
QIVGLNAMLMELSKNKNFSFINATSVPIIDNIESLTSNIRNTSSLGPYSTDAVKTVDLKLAGSGLMSKSSRR
FSGGILNRKAFEKYSLDTIVSYKNNDIRFYLILFGNDPINSKLQYLVNETGGAVIPFSSYEGVSKVYD
GTYLLEVVYPGPOEPNKYFNLSVEANINOOTGRGEFAYFIN

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AACTTGAAAAGTGCTTCTTTGATAATTCTATTATAAAAGGCCACATGGGTACCAAGATTACATCTTAAGGGC
TGATAAGTACGGGGATATTACGCTGCTAACATTGTACGCCATGAAATATTGTATTGATGTTAAACAATGTT

TABLE 1. Nucleotide and Amino Acid Sequences

AATGCTTTAGTTAAAGATGGCTTAGTTATTAACCTTATGATGTTATTGAAGCTAATAATCTGCTTTATG
 TGACTCTTATTCAAGTGTGAAATTGGTGTATGACAAAGTTCTGGAGTTAAAGGAAATCTATTGGAATAA
 AGGCACAAAAGATGGCGAATTGCTGCTCAGTATATGGCTATTGATAAGAGAAACTATATTGTAAGTGAG
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 AGTGATTTGAGGTAAATGTTAGTATCCTAAAAGATGGTGTATAAATATAGCATTGCTAAAAGACAA
 TTACAAAATTTAAAGCAGATAAAATGAATTCTAAAATTCTTCATCTATTGATGCCATAATCAGATGAT
 TGTCTCAGATTTAATAATGCCAAGGTTCAAGAGTGATGCAAGCCTTATGATAGTTAAATGTTGAT
 GTTAAAGAATAATTAGGCTTGGAGGGCTAAAATTACGTTAGCAGTAAAGCGGATTACAG
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 CAAATTGTAGGGTTAAATGCCCTAATGGAGTTGCAAAAAAATTTAGTTATAAATGCAACAAGTGTGC
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 TGTAAAACAGACGTTAGTTGAAGTTGCAGGTTCTGGCTTATGTCAAAAGCTCAAGAAGAGCAGTAGTTTAT
 TTTAGTGGTGGTATTTAAATCGTAAAGCTTTGAAAAGTACTCTTGTAGATAATAGTAAAGCTCAGTATTAGTAAAC
 ATGATATAAGGTTTACTTAATACTATTGTAATGATCCTATTAAAGTAAAGCTCAGTATTAGTAAATGAAAC
 AGGCGGTGCTGTAATTCTTTCATCTTATGAAGGTGTATCTAAAGTTATGATTAAATTAGAACAACAAAAGC
 GCCACTTATTGTTGGAATTATTATCCAGGCCCTAAGAACCTAATAAATATTAAATTATCTGTTGAAAGCAA
 ATATAAATCAACAGACAGGAAGAGGGAGTTGCATATTATTAAATTAG

f730.aa

MIKSILDYLLTLHPVLLGLGSTFTWFTTAFGAAAVFFRKVDNKIMDAMLGFSAGIMIAASFFSLIQPAIERAEE
 LGYITWVPAVFGFLVGAFFIYIVDVFPVPLDKLTFIDEDELTGHGKDFLLFTAVTLHNFPPEGLAVGVAFGALASNP
 DIQTLVGAMLLTLGIGIQNIPEGAAISLPLRRGNVALAKCFNYQMSGLVEIVGGLMGAYAVYSFTRILPFALAFS
 AGAMIVYVSIQLIPEAKRKDIDNKVPSIFGVIGFTLMMFLDVSLGZ

t730.aa

AVFFFRKVDNKIMDAMLGFSAGIMIAASFFSLIQPAIERAEEELGYITWVPAVFGFLVGAFFIYIVDVFPVPLDKLTF
 IDEDELTGHGKDFLLFTAVTLHNFPPEGLAVGVAFGALASNPDIQTLVGAMLLTLGIGIQNIPEGAAISLPLRRGN
 VALAKCFNYQMSGLVEIVGGLMGAYAVYSFTRILPFALAFSAGAMIVYVSIQLIPEAKRKDIDNKVPSIFGVIGFT
 TLMMFLDVSLGZ

f730.nt

ATGATAAAATCAATTAGATTATTAACCTTGCATCCTGTATTGGACTTTAGGTCTACTTCACCTT
 GGTTTACTACAGCTTTGGAGCAGCAGCAGCTTTCTTAGAAAGGTAGATAATAAAATGGACGCTATGCT
 TGGTTTCTAGCTGGCATTATGATAGCGGCCAGTTTCTCGCTTATTGAGCTATAGAAAGAGCTGAAGAG
 CTTGGATACATTACTGGGTGCCGGCTGTTGGATTCTGTTGGCATTCTTATATATTGAGATGTAT
 TTGTTCCAGATCTGGATAAAACTACTTTATTGATGAAGACTTAACAAACATGGAAAAAGATTCTTACTCTT
 TACTGCTGTTACTTACATAATTCCAGAAGGATTGGCTGTTGGAGTTGCTTTGGAGCCTGGCGTCAATCCA
 GATATTCAAACCTTAGTTGGGCTATGCTCTACGCTTGGTATTGGTATTCAAATATTCCCGAAGGAGCAGCTA
 TTTCTCTGCTTTAAGAAGAGGTATGTTGCTTGGCAAAATGCTTAACTATGCCAAATGTCAGGATTGGTAGA
 AATTGTTGGGGGGCTTATGGGTGCTTATGCCGTTATTCTTTACTGAATTTCACCTTGTGTTGGCTTTCT
 GCAGGAGCTATGATTATGTCATTGAAACAATTACCTGAAGCTAAGAGAAAAGACATTGACAATAAGTGC
 CAAGTATATTGGTGTATTGGTTACATTAATGATGTTCTCGATGTTCACTAGTTAA

t730.nt

GCAGTTTTCTTAGAAAGGTAGATAATAAAATGGACGCTATGCTTGGTTTCAGCTGGCATTATGATAG
 CGGCCAGTTTTCTGCTTATTCAGCCTGCTATAGAAAGAGCTGAAGAGCTGGATACATTACTGGGTGCCGGC
 TGTTTTGGATTCTGTTGGCATTCTTATATATATTGAGATGTATTGTTCTAGATCTGGATAAAACTTACT
 TTTATTGATGAAGACTTAACAAACATGGAAAAAGATTCTTACTGCTGTTACTTACATAATTTC
 CAGAAGGATTGGCTGTTGGAGTTGCTTTGGAGCCTGGCGTCAATCCAGATATTCAAACCTTAGTGGGCTAT

TABLE 1. Nucleotide and Amino Acid Sequences

GCTTCTTACGCTTGGTATTGGTATTCAAAATATTCCCGAAGGAGCAGCTATTCTCTGCCTTAAGAAGAGGTAAT
 GTTCTTGGCAAAATGCTTAACTATGCCAAATGTCAGGATTGGTAGAAATTGTGGGGGGCTATGGGTGCTT
 ATGCGGTTATTCTTACTCGAATTTCACCTTCTGCTTGGCTTCTGCAGGAGCTATGATTATGTCAT
 TGAACAATTAATACCTGAAGCTAAGAGAAAAGACATTGACAATAAGTCCAAGTATATTGGTGTATTGGTTT
 ACATTAATGATGTTCTCGATGTTCACTAGGTTAA

f197.aa

MLLKLKYRFGFLFLIFILLFSTIFNFVLCGYLEDYYKQLTRAQVRRAFSLQSFLDTLHVIINGAASNLALE
 TISEFAMSENRGKDFSESELIDLKNPKFVIDSVKSKYRQYLYNFMANLKNDTLFEFAFFDFEGRVIVSTRHE
 NNMDFGHSEANTNYFKKAVEDYRQNQLKFIGWYSLNSEGISAEVAIRSKQSEKKAFIAIVPVYSPEDKLVCGYLAG
 YLLNDIVADSFDRFRFGFYKRGNFIFYVDPNNIAVNPFEENETSRSVSSKFLNVLKDVFSKPPPSNIASESVYTI
 DRILLSEMGEDCYYAMLIPISSKLGEKSGVLIARLPYKDIYGVISLRFQYILYSVLGIIALSIVLSIRIDRIISFR
 LNAIRVLVQDMVKGNLDKDYALDDDENTLDEGMLSLOQVVKMKAISVAISSVLRNISYVNKASLEVASSSQNLSS
 SALQQASALEEMSANVEQIASGVNMSANNSYETEQIAALKTNENSQIGGRAVEESVIAMQDIVEKVSIEEIARKTN
 LLALNAAIEAARAGDEKGKFAVVASEIRKLADLSKISALEIGELVEDNSKVATEAGVIFKEMLPEIEETANLVKKI
 SEGSSKQSDQIAQFKMALDQVGEVVQSSASSSEQLSSMSDKMLEKSKELRKSVLFFKIKDSKIEPNENDDYDFRLI
 DCPENSKDENQNLKSNGISTSNASGHNNYSLDIESESSVRTINKRVDPKKAIIDIADKDLNFDDDFSEF

t197.aa

VLCGYLEDYYKQLTRAQVRRAFSLQSFLDTLHVIINGAASNLALETISEFAMSENRGKDFSESELIDLKNPKFV
 IDSVKVSKYRQYLYNFMANLKNDTLFEFAFFDFEGRVIVSTRHENNNMDFGHSEANTNYFKKAVEDYRQNQLKFI
 GWYSNLSEGISAEVAIRSKQSEKKAFIAIVPVYSPEDKLVCGYLAGYLLNDIVADSFDRFRFGFYKRGNFIFYVDPN
 NIAVNPFEENETSRSVSSKFLNVLKDVFSKPPPSNIASESVSVYTIDRILLSEMGEDCYYAMLIPISSKLGEKSGV
 IARLPYKDIYGVISLRFQYILYSVLGIIALSIVLSIRIDRIISFRNNAIRVLVQDMVKGNLDKDYALDDDENTLD
 ELGMLSLOQVVKMKAISVAISSVLRNISYVNKASLEVASSSQNLSSALQQASALEEMSANVEQIASGVNMSANNS
 YETEQIAALKTNENSQIGGRAVEESVIAMQDIVEKVSIEEIARKTNLLALNAAIEAARAGDEKGKFAVVASEIRKL
 ADLSKISALEIGELVEDNSKVATEAGVIFKEMLPEIEETANLVKKISEGSSKQSDQIAQFKMALDQVGEVVQSSAS
 SSEQLSSMSDKMLEKSKELRKSVLFFKIKDSKIEPNENDDYDFRLIDCPENSKDENQNLKSNGISTSNASGHNNY
 SLIDIESESSVRTINKRVDPKKAIIDIADKDLNFDDDFSEF

f197.nt

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 AGCTTTCTTGCAATCTTTAGACACCCCTGCATGTCATAATCAATGGTCAGCTCTAATTGGCAGTTGAA
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 AGTGCATTGCAACAGGCATCTGCTCTTGAAGAAATGTCAGCTAATGTTGAGCAAATAGCCTCAGGTGTCAACATGA
 GCGCCAATAATTCTTATGAAACAGAACAAATAGCTTAAAGACGAATGAAATTCTCAGATAGGTGGTAGGGCCGT
 TGAAGAATCTGTTATTGCTATGCAAGACATTGAGGAAAGTTAGTGTATTGAAGAGATAGCTAGAAAACCAAT
 TTACTGCTTGAATGCGGCTATTGAAGCTGCAAGAGCAGGAGATGAGGAAAGGGATTGCTTGTGCCAGTG

TABLE 1. Nucleotide and Amino Acid Sequences

AGATTAGAAAGTTGGCTGATTGAGTAAAATTCTGCTCTGAGATTGGAGAGTTAGTTGAAGATAACTCTAAGGT
 AGCAACTGAAGCGGGAGTGATCTTAAAGAAATGCTACCGAAATTGAAGAAACGGCTAATCTTGTAAAGAAGATT
 TCAGAAGGTAGCTCTAAGCAAAGCGATCAGATTGCTCAATTAAAATGGCTTAGATCAGGTTGGAGAAGTTGTC
 AATCTTCAGCTCAAGCAGTGAGCAGCTTAGTATGTCGATAAAATGTTAGAAAAGTCTAAGGAACTTAGAAA
 ATCTGTATTATTTTCAAAATTAAAGATTCTAAAATTGAAAATCCAGAAAATGATGATTATGATTCAGGTTAATA
 GATTGTCCTGAAAATTCTTTAAAGATGAAAATCAAATTGAAAAGCAATGGAATTCTACTTCAAATGCCAGTG
 CGCATAATAATTATCTTTAGATATTGAGAGCGAATCTCTGTAAGAACTATTAAAGCGAGTTGATCCTAAAAAA
 AGCTATCGATATTGCTGATAAGGATTAAATTGATGATGATTTTCAGAGTTTAG

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GTTTATGCGGTTATTAGAAGATTATTATAAGCAGCTACAAGGGCGCAAGTAAGAAGAGCAGCTTTCTTGC
 AATCTTTTAGACACCCCTGCATGTCTAATCAATGGTCAGCTCTAATTGGCACTTGAAACCATATCAGAATT
 TCGAATGTCTGAGAATAGAGGAAAAGATTCTCTGAGTCGGAATTGATAGATTAAAGAAAAAATCCAAAATTGTT
 ATTGACTCTGAAAGGTGAGCAAAAAATATCGACAATACTTATAACAATTATGGCCAATCTAAAAATGATAACCC
 TTTTGAGAATTGCTTTTGATTGAGGGAGAGTAATTGTTAGCACAAGACATGAGAATAATATGGATT
 TGGTCATTCTGAGGCTAATACCAATTATTAAAAGCTGTTAGGATTATAGGAAAACCAATTAAAATTTATA
 GGGTGGTATTCAAATCTTCTGAGGAATATCCGAGAAGTTGCTATTAGGTCTAAACAAAGCGAAAAAGGCTT
 TTGCAATAATTGTACCTGTATATTCCCAGAAGATAAACTGTTGTTGGGATTGGCCGGATATTGCTTAATGA
 TATTGTCAGATAGTTGATAGATTAGTCGTTTATAAAAGAGGCAATTATTATGTGGATCCAAC
 AATATAGCAGTTAACCTTTGAGAATATAATGAAACAGCAGGGTTAGTTCTAAATTGAAATGTTCTAAAG
 ATGTTTCTCTAACGCCCTTCCATCAAACATTGCCAGTGAAGTGTGGTTACACTATTGATAGAATACTTT
 GTCCGAAATGGGAGAAGATTGTTATTGCAATGTTGCCATAAGTAGTAAATTGGGAGAAAAGAGTGGAGTACTT
 ATTGCTAGGCTCCTTATAAGGATATTACGGAGTAATATCTAGTCTAAGATTCTAGTCTAAATTGCTTAATGA
 TAGGCATTATAGCATTAAAGTATTGTTCTTCATTAGAATAGACAGGATTATTGCTTAAACGCAATTAG
 AGTTCTAGTCAAGATATGTTAAGGGCAATTAGATAAGATTGCTCTGATGATGAAATACTCTGAT
 GAACCTGGCATGTTAAGTCTCAGGTTAAAATGAAAAAGCTATTCTGTAGCAATTCTAGTGTGAGAA
 ATATTAGCTATGAAATAAGGCAAGTTAGAAGTTGCCAGTCAAGTCAAATTAAAGCTCTAGTGCATTGCAACA
 GCCATCTGCTCTGAAAGAAATGTCAGCTAATGTTGAGCAAATGCCAGGTGTCAACATGAGCGCAATAATTCT
 TATGAAACAGAACAAATAGCTTAAAGACGAATGAAAATTCTCAGATAGGTGGTAGGGCGTTGAAGAATCTGTTA
 TTGCTATGCAAGACATTGGAGAAAGTTAGTGTATTGAGAGATAGCTAGAAAACCAATTACTGCTTGAA
 TGCGGCTATTGAGCTGCAAGAGCAGGAGATGAGGGAAAGGGATTGCTGTTGGCCAGTGAGATTAGAAAGTTG
 GCTGATTGAGTAAATTCTGCTTGTGAGATTGGAGAGTTGAGATAACTCTAAGGTAGCAACTGAAGCGG
 GAGTGTATTAAAGAAATGCTACCGAAATTGAAGAAAACGGCTAATCTTGTAAAGAAGATTGAGAAGGTTGCT
 TAAGCAAAGCGATCAGATTGCTCAATTAAAATGGCTTAGATCAGGTTGGAGAAGTGTCAATTCTCAGCTTCA
 ACCAGTGGAGCAGCTCTAGTATGTCGATAAAATGTTAGAAAAGTCTAAGGAACTTAGAAAATCTGTTATT
 TCAAAATTAAAGATTCTAAAATTGAAAATCCAGAAAATGATGATTATGATTTCTAGGTTAATAGATTGCTGAAA
 TTCTTTAAAGATGAAAATCAAATTGAAAAGCAATGGAATTCTACTTCAAATGCCAGTGGGCATAATAATTAT
 TCTTAGATATTGAGAGCGAATCTCTGTAAGAACTATTAAAGCGAGTTGATCCTAAAAAGCTATCGATATTG
 CTGATAAGGATTAAATTGATGATGATTTTCAGAGTTTAG

f200.aa

MTISKNVFSKFILKFLNSSAFVSFALFVGFLIVGLVVMGLGHSPFRMYFILEIIFSSPKHLGYVLSYSAPLIFT
 GLSIGISLKGALFNIGVEGQFILGSIVALIASVLLDLPILHVITIFIITFLASGSLGILIGYLKAKFNISEVISG
 IMFNWILFHNNIILDFSFIRDNDFSKPIKESAYIDFLASWKLSPLEGAYRSSHPFVNELLKAPLHFGIILGII
 FAILIWFLNKTIIIGFKINATGSNIEASRCMGINVKAVLIFSMFLSAAVAGLAGAIQLMGVNKAIFKLSYMQGIGF
 NGIAASLMGNNSPIGIFSSILFSILLYGSSRVQSLMGLPSSIVSLMMGIIVLVISASYFLNKIVLKGVKRVKYN
 ILD

t200.aa

LVVMGLGHSPFRMYFILEIIFSSPKHLGYVLSYSAPLIFTGLSIGISLKGALFNIGVEGQFILGSIVALIASVLL
 DLPPILHVITIFIITFLASGSLGILIGYLKAKFNISEVISGIMFNWILFHNNIILDFSFIRDNDFSKPIKESA
 YIDFLASWKLSPLEGAYRSSHPFVNELLKAPLHFGIILGIIFAILIWFLNKTIIIGFKINATGSNIEASRCMGINV

TABLE 1. Nucleotide and Amino Acid Sequences

KAVALIFSMFLSAAVAGLAGAIQLMGVNKAIFKLSYMQGIGFNGIAASLMGNNSPIGIIFSSILFSILLYGSSRVQS
LMGLPSSIVSLMMGIIVLVISASYFLNKIVLKGVKRVKYNNILD

f200.nt

ATGACAATTAGTAAAAACGTATTTAGTAAATTATTTGAAATTAAATTCTCAGCATTGTTAGGTATTTG
CTCTATTGTTGGATTAAATTGTTGGCTAGTGGTATGGGCTTGGCATTCTCCTTTAGAATGTATTTAT
AATATTAGAATTATTTCTCCAAACATTAGGTTATGTTAAGTTATTCAAGCTCCTTGATTTTACA
GGTCTTCATTGGTATTCTTAAAGCGGGCTTTAATATTGGGTTGAAGGCCAGTTACTAGGATCTA
TTGTTGCTTAATAGCATCAGTTACTTGATTTGCCAATTACATGTAATTACTATTTTATTACTTT
TTAGCTTCAGGCAAGTTAGGAATTAAATCGGATATTAAAGCCAATTCAATATTAGCGAAGTGAATTCA
ATAATGTTAATTGGATATTTCATTAAATAATATAATTAGTTAGTTATTAAAGAGATAATAGTG
ATTTCAAAACCCATTAAAGAAAGCGCATATTGATTAGCTTCTGGAAAGCTCTCACAGAAGGTCTGC
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TTGCTATTAAATATGGTTTACTTAATAAACTATTATTGATTTAAATAGCCACAGGAAGTAATATTG
AAGCTTCAGATGTATGGTATTAATGTAAGCTGTGTAATTTCATGTTCTCAGCAGCTGGCAGG
TCTTGCTGGTGTATTCAACTTATGGGTTAATAAGCTATTAAAGCTTATATGCAAGGAATTGGTTT
AATGGGATAGCTGCTCTTATGGGAAACAATTGCCAATTGGCATAATTAGCTTCTATATGCAAGGAATTGG
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TGTTCTGTAATTCTGCTAGCTATTAAATAAAATTGTTAAAGGTGTTAAGCGTGTCAAATAATAATT
ATTCTTGATTAG

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GGGCTAGTGGTATGGGCTTGGCATTCTCCTTTAGAATGTATTTATAATTAGAAATTATTTCTTCTC
CCAAACATTAGGTTATGTTAAGTTATTCAAGCTCCTTGTATTACAGGTCTTCTATTGGTATTCTTAA
AGCGGGCTTTAAATATTGGGTTGAAGGCCAGTTACTAGGATCTATTGTTGCTTAATAGCATCAGTTA
CTTGATTTGCCAATTACATGTAATTACTATTATTACTTTAGCTTCAGGAGTTAGGAATT
TAATCGGATATTAAAGCCAATTCAATATTAGCGAAGTGAATTCAAGGAATAATTGTTAATTGGATATTCTCA
TTAAATAATATAATTAGTTAGTTATTAAAGAGATAATTAGTGAATTTCAAAACCCATTAAAGAAAGC
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ATGAGCTTAAAGCACCTCTCATTTGGAATAATTAGGTATAATTGCTATTAAATGGTTTACT
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GTGTTAATAAGCTATTAAAGCTTCTTATGCAAGGAATTGGTTAATGGGATAGCTGCTCTTATGG
AAACAATTGCCAATTGGCATAATTAGCTTCTAGCATTCTTCTATATTGCTTATGGAAGCAGTAGAGTTCAA
AGTTTAATGGCCTTCATCTCAATTGATCTTGATGGAATAATTGTTCTGTAATTCTGCTAGCTATT
TTTAAATAAAATTGTTAAAGGTGTTAAGCGTGTCAAATAATAATTCTTGATTAG

f208.aa

MVKKFSIFLKAIIFSIPELLIEELSIILFLPYKIRFALIFLGFLFDTIFIFIFLYKITKAYLSQRLEIYVRNNLF
FDIIHCLIPLAFYSSYQLKNIIVAHETILNPIMLSFKLRFRLRRFNDLIIIEIYNSKEKNLILIAFARTFSMSL
LIPFTFFIIISSSKIVNSIPEKQEFNIIKNISIINEKAYIKEKYPFILIIKEKDDIIYSKSDEIFVYSPSEYRVI
EMEKTKFYIDKYLQRKSDSILGIFLFTLFASFTIFLMNFYKFFKASFLNPIILMTKILQDPLEYRKIQIPFTLSEE
KVYELAKSFNNLLKEKLNKRKSPIPLEIEVKKIINKNQEIK

t208.aa

IIIFSIFELLIEELSIILFLPYKIRFALIFLGFLFDTIFIFIFLYKITKAYLSQRLEIYVRNNLF
FYSSYQLKNIIVAHETILNPIMLSFKLRFRLRRFNDLIIIEIYNSKEKNLILIAFARTFSMSLIPFTFFIIIS
SSKIVNSIPEKQEFNIIKNISIINEKAYIKEKYPFILIIKEKDDIIYSKSDEIFVYSPSEYRVI
EMEKTKFYIDKYLQRKSDSILGIFLFTLFASFTIFLMNFYKFFKASFLNPIILMTKILQDPLEYRKIQIPFTLSEE
KVYELAKSFNNLLKEKLNKRKSPIPLEIEVKKIINKNQEIK

f208.nt

TABLE 1. Nucleotide and Amino Acid Sequences

ATGGTAAAAAAATTTCAATTCTTAAAGCAATAATAATTTCATATTGAACCTTTAATCGAAGAACTCT
 CAATAATTCTTTTACCATACAAAATACGATTCGACTAATATTCTGGGTTCTATTGACACAATTTCAT
 TTTCATTTTATACAAAATACCAAGGCCACCTTCCAAAGATTAGAAATCTACGTCAAGAAACAATCTATT
 TTCGATATAATCCACTGCCTTATTCTTAGCGTTATAGCTCATATCAGCTAAACATAATTGTCGCCATG
 AAACAATATTAAATCCAATAATGCTATCACTTTCAAGTTAAGATTTCAGCTTAAATGACCTAAT
 AATAGAAATATATTACAATTCAAAAGAAACCTAATACTAATAGCATTGCTAGGACATTTCATGAGCTTA
 TTAATACCATTTACATTTTATAATAATCAAGCTCAAAATTGTAATTCAATACCAGAAAACAAGAATT
 ATATCATTAAAATATCAATAATAATGAAAAGCTTACATTAAAGAAAATATCCCTCATCTTAATAATCAA
 GGAAAAGAGTGAACATAATACAGACGAAATTGTTACTACAGTCCAGTGAATATAGAGTAATA
 GAAATGGAGAAAACAAAATTTATATAGATAAAATTGCAAGAAAAGCATTCTATTCTGGAAATTTCAT
 TTACATTGTTGCATCATTTACTATTTTAATGAATTTCATTTAAAGCAAGCTTTAAATCCTAT
 TATTTAATGACAAAATTTCAGACCCATTAGAATATCGAAAATTCAAATTCTTACTTTAAGCGAAGAA
 AAAGTATATGAACTGCAAAATCATTAACAATCTCTGCTAAAGAAAACCTAAACTCAAAGCGAAAAGCAA
 TACCTTAGAAATTGAAAAGTAAAAAAATAATTAAATAAAACCAGGAAATAATGA

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ATAATAATTTCATATTGAACCTTTAATCGAAGAACTCTCAATAATTCTTTTACCATACAAAATACGAT
 TTGCACTAATATTCTGGGTTCTATTGACACAATTTCATTTTATACAAAATAACCAAGGCCA
 CCTTTCCAAAGATTAGAAATCTACGTCAAGAAACAATCTATTCTCGATATAATCCACTGCCTTATTCTTAGCG
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 TCAAGTTAAGATTTCAGACTCTTAGGTTAATGACCTAATAATAGAAATATTTACAATTCAAAGAAAAGAA
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 AGCTCAAAATTGTAATTCAATACAGAAAACAAGAATTAAATATCATTAAATATCAATAATAATGAAA
 AAGCTTACATTAAAGAAAATATCCCTCATTTAAATCAAGGAAAAGATGACATAATATACTCAAATCAGA
 CGAAATATTGTTACTACAGTCCAGTGAATATAGAGTAATAGAAATGGAGAAAACAAAATTTATATAGATAAA
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 TGAATTTCATAAATTTCAGCAAGCTTTAAATCCTATTAAATGACAAAATTTCACAAGACCCATT
 AGAATATGAAACATTCAAATTCTTACTTTAAGCGAAGAAAAGTATATGAACTGCAAACATTAAACAT
 CTCTGCTAAAGAAAACCTAAACTCAAAGCGAAAAGCAAACCTTAGAAATTGAAAAGTAAAAAAATAA
 TTAATAAAACCAGGAAATAATGA

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MKIQIIIMLLALLDPLNARLLDISIEKRADEEIKKYSSYNLILEKEYTNFPTSEIEKNIYKLTEHFVKSIMLNK
 TNYSLLNSNYKEANKYLIQSELIDKKFLKYKIFKIKNINGIFKSHSLIYTKKGFYKLELYIENNAEPLKIFNLNIT
 YFLKNLDKISNEMIFFPREKREVNMIQKTTIAADSSKPRGINYDTGIPFNVLIVDDSVFTVKQLTQIFTSEGPN
 IDTAADGEEAVIKYKNHYPNIDIVTLDITMPKMDGITCLSNIMEFDKNARVIMISALGKEQLVKDCLIKGAKTFIV
 KPLDRAKVLQRVMSVFVK

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RLLDISIEKRADEEIKKYSSYNLILEKEYTNFPTSEIEKNIYKLTEHFVKSIMLNKTNYSLLNSNYKEANKYLIQ
 SELIDKKFLKYKIFKIKNINGIFKSHSLIYTKKGFYKLELYIENNAEPLKIFNLNITYFLKNLDKISNEMIFFPRE
 KREVNMIQKTTIAADSSKPRGINYDTGIPFNVLIVDDSVFTVKQLTQIFTSEGPNIDTAADGEEAVIKYKNHYP
 NIDIVTLDITMPKMDGITCLSNIMEFDKNARVIMISALGKEQLVKDCLIKGAKTFIVKPLDRAKVLQRVMSVFVK

f210.nt

ATGAAAATTCAAATAATTATAATGCTGCTGCATTGTTAGATTTCACCTTAATGCCAGACTTTGGACATTCAA
 TTGAAAAGAGCAGATGAAGAAATAAAAATATTCTGTCTTATAATTAAATTGAAAAGAAATCTACCAA
 TTTTCCAAACAGCGAAATAGAAAATATTATAAACTACAGAACATTGTAAAAAGCATAATGCTCAATAAA
 ACTAACTACAGCTATTAAATTCAAACCTACAAAGAACAAATAATCTAATTCAAAGCGAACTCATTGATAAAA
 AATTTCATAAATATAAAATATTAAATGAAATTTCAGCATTCACTAATATACAAA

TABLE 1. Nucleotide and Amino Acid Sequences

AAAAGGATTTACAAATTAGAACCTTACATAGAAAATAATGCAGAACCTCTAAAATATTAACCTAACATTACT
TATTTTTAAAGAATTAGATAAAATAAGTAATGAAATGATTTTCCCAAGGGAATGA

t210.nt

AGACTTTGGACATTCAATTGAAAAAGAGCAGATGAAGAAATAAAAATATCGTCTTATAATTAAAGTAG
AAAAGAACATACTATACCAATTTCACAAGCGAAATAGAAAAAATATTATAAAACTAACAGAACATTGTAAA
AAGCATAATGCTCAATAAAACTAACAGCTTATTAAATTCAAACATACAAAGAACATAATCTAATTCAA
ACCGAACATGATAAAAATTTAAAATATAAAATTTAAAATCAAACATATAAATGAAATTAAAGGCC
ATTCACATAATATAACAAAAAGGATTTACAAATTAGAACCTTACATAGAAAATAATGCAGAACCTCTAAAAT
ATTTAACCTAACATTACTTATTTTAAAGAATTAGATAAAATAAGTAATGAAATGATTTTCCCAAGGGA
TGA

f22.aa

MLKTLTKIITISCLIVGCASLPYTPPKQNLNYLMELLPGANLYAHVNLIKNSIYNNSLPKYKSVLGLISNLYFSY
KKENNDFAILLIMGNFPKDFIFWGHIKRNNTESIGNIFTNPWKLKNSNIYIIPNKARTSIAITQKDITAKDNNMLTT
KYIGEIEKNEMFFWIQDPTLLPNQIVSSKNLIPFSSGTLINSLNQEEYIFKSLIKTNNPPILKILSKKLIPTVL
TNMTNLTISSHIKTTIKDQNTVEIEFNIQKSSVESLIEKLASNIQT

t22.aa

PYTPPKQNLNYLMELLPGANLYAHVNLIKNSIYNNSLPKYKSVLGLISNLYFSYKKENNDFAILLIMGNFPKDFIW
GIHKRNNTESIGNIFTNPWKLKNSNIYIIPNKARTSIAITQKDITAKDNNMLTTKYIGEIEKNEMFFWIQDPTLL
LPNQIVSSKNLIPFSSGTLINSLNQEEYIFKSLIKTNNPPILKILSKKLIPTVLTNMTNLTISSHIKTTIKDQNT
VEIEFNIQKSSVESLIEKLASNIQT

f22.nt

ATGTTAAAACATTAACAAAATAATTACCATTTCATGCCTCATAGGGATGCGCAAGCCTGCCCTACACTCCTC
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CAGGTCTATTATAACTCTTAAGCCCTAAATATAAACTAGTTCTGGGTTATAAGCAATTATACTTAGCTAT
AAAAAGAAAATAACGATTTGCTACTAATAATGGTAATTCCAAAAGATATTCTGGGAATTCTATAAAA
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TCCAAACAAAGCTAGAACCTAGCATTGCAATAACCCAAAAGATATAACCGAAAAGACAATAATGCTAACAA
AAATATATTGGGAATAGAAAAATGAAATGTTTTGGATTCAAGATCCAACATTATTGCTCCAAACAAA
TAGTAAGCAGAAAATTAAATCCCTTAGCAGTGGAACTTGTCTATAACAGCTTAAATCAAGAAGAATATAT
TTTAAATCCTTAATCAAACAAATAATCCACCAATACTAAAATATTGCAAAAAGTTAATCCAACCGCTCTG
ACAAACATGACAAACCTCACAAATATCAAGCCACATAAAGACCACAATAAAGACCAAAACGTTGAAATAGAAT
TTAATATTCAAATCTAGTGTGAAAGCCTTATAGAAAAACTAGCTTCAAATATTCAAACCTAA

t22.nt

CCCTACACTCCTCCAAAACAAAATCTAAATTACTTAATGGAACCTTTACCTGGCGAAATTATACGCCATGTAA
ATTTAATTAAAACAGGTCTATTATAACTCTTAAGCCCTAAATATAAACTAGTTCTGGGCTTATAAGCAATT
ATACTTAGCTATAAAAAGAAAATAACGATTTGCTACTAATAATGGTAATTCCAAAAGATATTCTGG
GGAATTCTATAAAAATAGAAATACAGAACATAGCAATATTTACAATCCAAAATGGAAACTTAAAATTCAA
ATATATACATTATTCTAAACAAAGCTAGAACCTAGCATTGCAATAACCCAAAAGATATAACCGAAAAGACAATAA
TATGCTAACAAACAAATAATTGGGAAATAGAAAAAAATGAAATGTTTTGGATTCAAGATCCAACATTATG
CTCCCAAACCAAATAGTAAGCAGCAAAATTAAATTCCCTTAGCAGTGGAACTTGTCTATAACAGCTTAAATC
AAGAAGAATATATTAAATCTTAATCAAACAAATAATCCACCAATACTAAAATATTGCAAAAAGTTAAT
TCCAACCGTCTTGACAAACATGACAAACCTCACAAATATCAAGCCACATAAAGACCACAATAAAGACCAAAATACG
GTTGAAATAGAATTAAATTCAAACATGACAAACCTCACAAATATCAAGCCACATAAAGACCACAATAAAGACCAAAATACG
AA

f221.aa

TABLE 1. Nucleotide and Amino Acid Sequences

MGITVYLFISIFASFVLGSSMDSVKENVLKSTIFYDVEEVEFPYARKQTLQFIAKTHLKYAVFNFDKNKMF SYTF
VFDKKLISQYAIFIEVKKFGEATLVTPLNYLWDLGDSIIVLNKNILRITLKS YISNYNK

t221.aa

SMDSVKENVLKSTIFYDVEEVEFPYARKQTLQFIAKTHLKYAVFNFDKNKMF SYTFVFDKKLISQYAIFIEVKKF
GEATLVTPLNYLWDLGDSIIVLNKNILRITLKS YISNYNK

f221.nt

ATGGGTATTACAGTTTTTATTCTTTGATCTTTGTTCTGGGTTCTAGCATGGATTCTGTTAAAG
AGAATGTTCTCAAGAGCACTATTTTATTATGATGTTGAAGAAGTTGAATTTCCTATGCTAGGAAGCAGACTTT
ACAATTATGCTAAAACCCATTAAAATATGCTGTTTAATTGACAAAAATAAAATGTTCGTACACTTT
GTTTTGATAAAAATAATATCTCAGTATGCAATTTTATTGAGGTAAAGAAAAAGTTGGCGAGGCTACACTAG
TAACGCCTTGAATTATTGAGGTCTGGTATTCTATTGTTAAATAAAATTTAAGAATTACTTT
AAAATCTTATATTCAAATTATAATAATGA

t221.nt

AGCATGGATTCTGTTAAAGAGAATGTTCTCAAGAGCACTATTTTATTATGATGTTGAAGAAGTTGAATTTCCTT
ATGCTAGGAAGCAGACTTACAATTATTGCTAAAACCCATTAAAATATGCTGTTTAATTGACAAAAATAA
AATGTTTCGTACACTTTGTTGATAAAAATAATATCTCAGTATGCAATTTTATTGAGGTAAAGAAAAAG
TTTGGCGAGGCTACACTAGTAACGCCTTGAATTATTGAGGTCTGGTATTCTATTGTTAAATAAA
ATATTGTTAAGAATTACTTAAATCTTATATTCAAATTATAATAATGA

f253.aa

MYMENIEVRGQPNNFFGLIPFFVIIYLGTGIYLGIVIGVEMAFYQLPASVAMFFASIVCFLVKFKFSDKIHF IK
GAAQYDIILMCLIFMLSGAFSSLCKEIGCVETVANLGKYINPNWIVSGIFFVTCFLSFSAGTSVGSIVAI APIAF
NIAVKSGINPNLIAASVMCGAMFGDNLSLISDTTIVSSRTQGSSILDVFISSSFYAFPSAILTFFSFFFSENLSN
ATNFLHESSIDLVKTVPYLMIIFFSLAGMNVFIVLFLGILSICLISVLYGNLYFLDVMKNINKFLNMA DLIFLSI
LTGGVSFAVIHNGGFKWLLIKLKS LIRGKSSAEFSIGAFVSIVDVFLANN TIAIILICGVAKKIAFENNISVQRSA
SILDMFSCIFQGIIPYGAQMIILVNFNSNGLVSPISILPFLVYFGFLFFVILSILGLDIKKVFLFFLKK

t253.aa

LVFKFKFSDKIHF IKGAAQYDIILMCLIFMLSGAFSSLCKEIGCVETVANLGKYINPNWIVSGIFFVTCFLSFS
AGTSVGSIVAI APIAFNIAVKSGINPNLIAASVMCGAMFGDNLSLISDTTIVSSRTQGSSILDVFISSSFYAFPSA
ILTFFSFFFSENLSNATNFLHESSIDLVKTVPYLMIIFFSLAGMNVFIVLFLGILSICLISVLYGNLYFLDVMKN
INKFLNMA DLIFLSILTGGVSFAVIHNGGFKWLLIKLKS LIRGKSSAEFSIGAFVSIVDVFLANN TIAIILICGV
AKKIAFENNISVQRSA SILDMFSCIFQGIIPYGAQMIILVNFNSNGLVSPISILPFLVYFGFLFFVILSILGLDIK
KVFLFFLKK

f253.nt

ATGTATATGGAAAATATTGAAGTAAGAGGGCAGCCAAATTTTGGGCTTATTCCCTTTTTGTTATT
TCTATTAGGCACGGGATTATTGGGAGTTATTGGTAGAAATGCCCTTTATCAACTGCCGGCTAGTGGTGC
AATGTTTTGCTTCCATTGTTGGTATTAAAGAAAATTTCGACAAAATTCACTATTAA
GGAGCAGCTCAGTACGATATTACTAATGTGCTTATTGTTATGCTTCCGGAGCTTCTCTCTTTGTTAAAG
AAATAGGCTCGTTGAAACTGTAGCAAATTGGAAATTAAATATTAAATCCTAATTGGATTGTTCTGGTATATT
TTTGTAACCTGTTCTTCTTCTGCCGCACCTCTGGATCTCGTGCATTGCTCTATTGCTTT
AATATTGCTTAAAGCGGCATTAATCGAATTAAATAGCAGCATCTGTAATGTGAGCTATGTTGGAGATA
ATCTTCTTAAATATCAGATACAACATTGTTCTAGTCGAACCTCAAGGTAGTAGCATCTTAGATGTTTATTAG
TAGCAGTTTATGCTTCCATCGCCACTAACCTTTCTTCTTCTTGAAATTGCCAAT
GCCACAAACTTTACACGAAAGTTCAATAGATTAGTGAAGACTGTGCCTTATTAAATGATTATTTCTCTT

TABLE 1. Nucleotide and Amino Acid Sequences

TAGCTGGAATGAATGTTTTATAGTCTTTAGGTATTCTTCTATATGCTTATTAGCGTTTGTATGGTAA
 TTATACCTCTAGATGTAATGAAAAACATTAATAAAGGGTTTTAAATATGGCGATTGATTTCTTCATT
 TAAACAGGGGAGTTCTTCCGCTGATCATAATGGAGGTTAAATGGCTACTTATTAAATTAAACCTTGA
 TTAGAGGAAAAGTTCAAGCGGAATTCTATTGGGCTTTCAATAGTGTGTTCTGCTAATAACAC
 AATTGCCATACCTATTGCGGAAAGTAGCAAAAAGATAGCTTGAAGAAACATCAGTGTCAAAGAAGTGCT
 TCTATTAGATATGTTCTTGTATTTCAGGCATTATTCTTATGGTGCCTAATGATTATTTAGTGAATT
 TTCAAAATGGACTTGTGCGCAATTAGTATTGCAATTAGTTAGTTATTTGGATTTTATGTTTTGTTAT
 TTATCTATTGGCCTGATATAAAGTTTTTATTTTAAAGAATAA

t253.nt

TTGGTATTAAAGGAAAATTCCGACAAATTACATATTAAAGGAGCAGCTCAGTACGATATTACTAA
 TGTGCTTATTCTTATGCTTCGGGAGCTTCTCTCTTGTAAAGAAATAGGCTGCGTTGAAACTGTAGCAA
 TTGGGAATTAAATATTAATCCTAATTGGATTGTTCTGGTATATTGTTGTAACCTGCTTCTTCTTTCT
 GCCGGCACTCTGTTGGATCTCGTGCATTGCTCTATTGCTTTAAATTGCTGTTAAAGCGGCATTAATC
 CGAATTAAATAGCAGCATCTGTAATGTTGGAGCTATGTTGGAGATAATCTTCTTAAATACAGATAACATAT
 TGTCTAGTCGAACCTCAAGGTAGTAGCATCTTAGATGTTTATTAGTAGCAGTTTATGCTTCCATCCGCC
 ATACTAACTTTTTCTTCTTCTGAAATTGCTCAATGCCACAAACTTTACACGAAAGTCAA
 TAGATTTAGTGAAGAAACTGTCCTTAAATGATTATTTCTCTTAGCTGGAATGAATGTTTATAGTCT
 TTTTAGTATTCTTCTATATGCTTATTAGCTTGTATGGTAAATTACTTCTAGATGTAATGAAAC
 ATTAATAAAGGGTTTAAATATGGCGATTGATTTCATTCAATTAAACAGGGGAGTTCTTGGCCTGAA
 TTCATAATGGAGGCTTAAATGGCTACTTATTAAATTAAATCCTGATTAGAGGAAAAGTTCAAGCGGAATTTC
 TATTGGGCTTTGTTCAATAGTGTGTTCTGCTAATAACACAATTGCCACTTATTGCGGCAAAGTA
 GCAAAAAGATAGCTTGTAAACATCAGTGTCAAAGAAGTGCTCTATTAGATATGTTCTTGTATTT
 TTCAGGCATTATTCTTATGGTGCCTAATGATTATTAGTGAATTTCAAATGGACTTGTGCGCAATTAG
 TATTGCAATTAGTTATTGATTGTTATTGTTATTGTTATCTATTGTTGATATAAAAGTTTTTAAAGAATAA
 AAAGTTTTTATTTTAAAGAATAA

f265.aa

MRKCFVSLSLLIFFACSSNVEIELNDDISGIVSIFVNVRREFEKIRKELLTLVGEIANMPLFPVDEIKKYFKN
 GEEKLGLKLLSIKTQGDSINLVVKFDNLKILGDMKKPDISVFKIEKKDGKNIIELNINLENATKNINENKEYIS
 DALAALLPSDEIPMSAKEYKDVLVYFLSDFTSKASELIDNSKLNLVVKTSRNVQEFGFKQINSNLRFEMDMVKG
 LSLETPIKLRLV
 Y

t265.aa

SNVEIELNDDISGIVSIFVNVRREFEKIRKELLTLVGEIANMPLFPVDEIKKYFKNGEEKLGLKLLSIKTQGDS
 INLVVKFDNLKILGDMKKPDISVFKIEKKDGKNIIELNINLENATKNINENKEYISDALAALLPSDEIPMSAKE
 YKDVLVYFLSDFTSKASELIDNSKLNLVVKTSRNVQEFGFKQINSNLRFEMDMVKGSLLETPIKLRLVY

f265.nt

ATGAGAAAGTTTGTTAGCTTGTAGTTATTGTTGATTGTTGCTTGTAGCTCTAATGTTGAAATTGAGTTAA
 ATGATGATATTAGTGTATTGTTCAATATTGTTAATGTTAATAGAGAAATTGAAAGAAACTCTT
 AACAACTTGGTGGAGAAGAAATTGCAAAATATGCCCTTTCTGTAGATGAAATAAAACTTTAAAT
 GGAGAGGAAAAGCTGGCTTAAGCTTGTAGTATTAAACCAAGGAGATTCTATTAAATTAGTTGTTAAGTTG
 ATAATTAAATTAAATTAGGCGATTATGAAAAACCGATATCTGTGTTAAGATGAAAGATGG
 TAAATATTATTGAACTTAATATTAAATTGAAAACGCTACTAAGAATATTAAATGAAATAAGAATATTAGT
 GATGCACTTGCTGCTTTGCCATGGATGAGATCCAATGCTGCCAAAGAATATAAGATGTTTGGTTATT
 TTTATCGGATTTACTTCAAAGCAAGTGAACCTATTGACAATTCAAACCTTAATCTGTAGTTAAGACTCTAG
 AAATGTTCAAGAACAAATTGGATTCAAACAAATTAAACTCAAACACACTGCCGTTGAGATGGATGGTAAAGGA
 TTAAGTCTGAAACACCAATAAAACTAGATTAGTTATTGA

t265.nt

TABLE 1. Nucleotide and Amino Acid Sequences

TCTAATGTTGAAATTGAGTTAAATGATGATATTAGTGGTATTGTTCAATATTGTTAATGTTAATAGAGAATTG
 AAAAATTAGAAAAGAACTCTAACAACTTGGTGGAGAAGAAATTGCAAATATGCCCTCTTCTGTAGATGA
 ATAATTTACTTAAAATGGAGAGGAAAGCTGGGCTTAAGCTTTGAGTATTAAACCAAGGAGATTCT
 ATTAATTAGTTGTTAAGTTGATAATTAAATTAAATTAGGCATTATGAAAAACCGATATATCTGTGT
 TTAAGATAGAAAAAGATGGTAAAATATTGAACTTAATATTAAATTGAAAACGCTACTAAGAATATTAA
 TGAAAATAAAGAATATTAGTGTGCACTTGTCTTGTGCCATGGATGAGATCCAATGTCGCCAAAGAA
 TATAAAGATGTTGGTTATTTCATGGATTACTTCAAAGCAAGTGAATTGACAATTCAAACCTTA
 ATCTTGAGTTAAGACTCTAGAAATGTTCAAGAACATTGGATTCAAACAAATTAACTCAAACACACTGCGGTT
 TGAGATGGATATGGTAAAGGATTAAGTCTGAAACACCAATAAAACTAGATTAGTTATTGA

f269.aa

MNIRKLLFCIFFMNISFLFAGDYKGLDFKIKFFNQSIYRVNSNVFIEVSLSNASESVLTLEIGDINSFGFDFDVT
 DTTNIKVKRPIEYVKKRSKNAIPVRNMSLRPNEKFVSVINLNQFVKFSKDGVYFVKGIFFPDIIDPSKKKESNII
 TLFLNDGFDENPGSIDLVNLSENNIDQDILKKKKLSPDEIVKYLKALQLGKKEKFLYLDIEGLLNDKGKAYLY
 KQKLSPIPNKNVVEYKEYLWNSNNSDISKAPNKFSSIETTYSDTSGKVIADLYFDDQFYISKRYTFFKKYDYY
 WIIYDYIVQNTGIKEK

t269.aa

GDYKGLDFKIKFFNQSIYRVNSNVFIEVSLSNASESVLTLEIGDINSFGFDFDVTDTNIKVKRPIEYVKKRSKNA
 AIPVRNMSLRPNEKFVSVINLNQFVKFSKDGVYFVKGIFFPDIIDPSKKKESNIIITLFLNDGFDENPGSIDLVNL
 ENNDIQDILKKKKLSPDEIVKYLKALQLGKKEKFLYLDIEGLLNDKGKAYLYKQKLSPIPNKNVVEYKEYLW
 NSNNSDISKAPNKFSSIETTYSDTSGKVIADLYFDDQFYISKRYTFFKKYDYYWIIYDYIVQNTGIKEK

f269.nt

ATGAATATTAGAAAATTGCTTTTGATCTTTTATGAATATTCTTTCTTTGTTGCGGGAGATTACAAGG
 GCCTTGATTAAATCAAGTTTAATCAATCTATTATCGTCATAGTAATGTTTATTGAAGTTCTCT
 TAGTAATGCGTCTGAGAGTGTAACTTAGAAATAGGCATATTAAATTCTTGCTTGTATTGATGTTACT
 GATACCACCAATATTAAAGTAAAAGACCTATTGAATATGTTAAAAGAGATCTAAAATGTTCAATTCTGTTA
 GAAATATGAGCTTGAGACCTAATGAAAATTCTGAGTTATTAACTTAAATCAATTGTTAAGTTAGTAAAGA
 TGGAGTTATTGTTAAGGTATTCTCCAGACATTTCAGATCCATCTAAGAAAAAGAATCCAATTATT
 ACGCTTTTGAAATGATGGTTGATGAAAATCCAGGTAGCATAGACCTGTTAATTGTCGAAAATATGATA
 TTCAAGATATCTGAAAAGAAAAATTATCTCCGATGAAATTGTTAAATATTGTTAAGGCATTGAGCTTGG
 GAAAAAGAAAAGTTCTTTATATCTTGATATTGAAGGTTGTTATTAAATGACAAGGGCAAGGCATACCTTAT
 AAGAAAAGTTATCACCTATTCCAATAAAATGTTAGTTGAAGAGTATAAGAATATTGTTAAATTCTAATAATT
 CGGATATTCAAAAGACCAATAAATTCTATTGAAACTACTTATTCTGATACTTCTGGCAAGGTGATTGC
 TGATTATATTGACGATGGCAATTATTCAAAAGATACTTCTCTTAAAAATATGATTATT
 TGGATAATTATGATTACATTGTTCAAAACTGGCATTAAAGGAAAAGTAA

t269.nt

GGAGATTACAAGGGCCTGATTAAATCAAGTTTTAATCAATCTATTATCGTCATAGTAATGTTTTA
 TTGAAGTTCTCTTAGTAATCGCTCTGAGAGTGTAACTTAGAAATAGGCATATTAAATTCTTGCTTGT
 TTTGATGTTACTGATACCACCAATTAAAGTAAAAGACCTATTGAATATGTTAAAAGAGATCTAAAATGTT
 GCAATTCTGTTAGAAATATGAGCTTGAGACCTAATGAAAATTCTGAGTTATTAACTTAAATCAATTGTTA
 AGTTAGTAAAGATGGAGTTATTGTTAAGGGTATTCTCCAGACATTTCAGATCCATCTAAGAAAAAGA
 ATCCAATTATTACGCTTTTGAAATGATGGTTGATGAAAATCCAGGTAGCATAGACCTGTTAATTGTC
 GAAAATAATGATATTCAAGATATTGAAAAGAAAAATTATCTCCGATGAAATTGTTAAATATTGTTAAAGG
 CATTGCAGCTTGGAAAAAGAAAAGTTCTTTATCTGATATTGAAGGTTGTTATTAAATGACAAGGGCAA
 GGCATACCTTATAAGCAAAGTTATCACCTATTCCAATAAAATGTTAGTTGAAGAGTATAAGAATATTGTT
 AATTCTAATAATTGAGATTCTAAAAGACCAATAAAATTCTATTGAAACTACTTATTCTGATACTTCTG
 GCAAGGTGATTGCTGATTATTTGACGATGGCAATTATTCAAAAGATACTTCTTCTTAAAAA
 ATATGATTATTGAGATTACATTGTTCAAAACTGGCATTAAAGGAAAAGTAA

TABLE 1. Nucleotide and Amino Acid Sequences

f29.aa

MNWLSFFYVLLFLLIFPFELQSNKENIENLIKHLHMLYDLTNNLSELETINKIKNFDLEQHYLLITKYYLKIKKY
KEANDFLKKINQKKIKNQKIKNNEIISLKLKRINEDNINEEEIKKILNNEKNIDVKIIYQIFSLIKFKNKLANKIKN
IILTNYPKSIYSYKIKRNE

t29.aa

NNKENIENLIKHLHMLYDLTNNLSELETINKIKNFDLEQHYLLITKYYLKIKKYKEANDFLKKINQKKIKNQKIKN
EIISLKLKRINEDNINEEEIKKILNNEKNIDVKIIYQIFSLIKFKNKLANKIKNIIILTNYPKSIYSYKIKRNE

f29.nt

ATGAACCTGGCTATCCTTTTTATGTTTATTATTTTATTAAATTTTCTTTGAATTACAGAGTAATAATAAG
AAAATATAGAAAATTAAATAAGCTACATATGCTTTATGATTTAACCAATAACCTGTCAAAAGAATTAGAAACAAT
AAATAAAATTAAAAATTGGACTTAGAACACATTATCTGCTAATTACAAAATATTATCTAAAAATAAAAAAAATAT
AAAGAAGCTAATGATTTTAAAAAAATAAACCAAAAAAGATCAAAAATCAAAAATAAAAACGAAATCATT
CGCTAAAATTAGAATAATGAAGATAATATTAAATGAAGAAGAAATCAAAAATTTAAATAACGAAAAAAATAT
AGATGTCAAATAATTATCAAATATTCACTTATAAAATTAAAAATAAAATAGCAAATAAAATTAAAAAC
ATAATACTAACAAACTATCCAAAAGCATTATTCTTATAAAATAAAAGAAATGAATAA

t29.nt

AATAATAAAAGAAAATATAGAAAATTAAATAAGCTACATATGCTTATGATTTAACCAATAACCTGTCAAAAGAAT
TAGAAACAATAAAATAAAATTAAAAATTGGACTTAGAACACATTATCTGCTAATTACAAAATATTATCTAAAAT
AAAAAAATATAAGAAGCTAATGATTTTAAAAAAATAAACCAAAAAAGATCAAAAATCAAAAATAAAAAC
GAAATCATTGCTAAAATTAGAATAATGAAGATAATATTAAATGAAGAAGAAATCAAAAATTTAAATAACG
AAAAAAATATAGATGCTAAAATAATTATCAAATATTCACTTATAAAATTAAAAATAAAATAGCAAATAA
ATTAAAAACATAACTAACAAACTATCCAAAAGCATTATTCTTATAAAATAAAAGAAATGAATAA

f290.aa

MNSIYVIGKLLLTLFLIFFPFCYNLFANLAEINKLSEYAKSIVLIDFTKRILYSSKKPNLVFP PASLTKIVTIYT
ALIEAEKRNIKLKSIVPISDSASYYNAPPNSLMFLEKGQIVNFEIILKGLSVSSGNDSSIAIAEFVVGNLNSFVN
LMNINVNLGLFNMFVPEPGYSENKITALDMAFFVKSIEKFKFMLNIHSLKYFIPKSRNLGTALSSKFLNLK
QRNANLLIYDYPYSDGIKTGYIKESGLNLVATAKKERRLIAVVLGVEKGINGFGEKMRSSIAKNLFYEGFNKYSK
FPLIVKLKEKVYNGTVDTVALFSKEPFYYILTKDEFDKINISYTVDKLVAPLSGDMVGRAMIFLENEKIGDVALF
SGKVKRLGFQGLYKSFINLFREY

t290.aa

VNLAEINKLSEYAKSIVLIDFTKRILYSSKKPNLVFP PASLTKIVTIYTALIEAEKRNIKLKSIVPISDSASYYNA
PPNSSLMFLEKGQIVNFEIILKGLSVSSGNDSSIAIAEFVVGNLNSFVNLMNINVNLGLFNMFVPEPGYSENK
ITALDMAFFVKSIEKFKFMLNIHSLKYFIPKSRNLGTALSSKFLNLKQRNANLLIYDYPYSDGIKTGYIKESGL
NLVATAKKERRLIAVVLGVEKGINGFGEKMRSSIAKNLFYEGFNKYSKPLIVKLKEKVYNGTVDTVALFSKEPF
YYILTKDEFDKINISYTVDKLVAPLSGDMVGRAMIFLENEKIGDVALFSGKVKRLGFQGLYKSFINLFREY

f290.nt

ATGAATAGTATCTATGTTATTGGAAATTGTTATTAACTTTATTAAATTCCCCGTTTGTATAATCTT
TTGCAGTTAATTAGCTGAGATTAATAAATTATCAGAGTATGCAAAGTCATAGTTAATAGATTTGATACTAA
GCGAATACCTTATTCTAAGAACGCCAATTGGTTTCTCCAGCATCTTACAAGATTGTTACAATTATACA
GCTTAATTGAAGCTGAAAGCGAAATATAAAATTAAAAAGCATAGTTCTATTAGCATTCTGCTTATATTATA
ATGCACCCCCAATTCTCTTGATGTTAGAAAAAGGTCAAATTGTTAATTTGAAGAGATTAAAAGGACT

TABLE 1. Nucleotide and Amino Acid Sequences

TTCACTTCTCGGTAAATGATTCTCTATTGCAATTGCTGAGTTGAGTAGGCAATTAAATAGCTTGTAAATTTAATGAAATATTAAATTTAATGCTTAAATATGCATTGTTGAAACCTTCTGGATATAGCAGCGAGAATAAAGATTACAGCACTAGATATGGCTTTTGTGAAATCTTATATAGAAAAGTTAAATTATGCTTAATATTCACTTCTTAAAGTATTATTTATCAGGAAAGTAGAAATTAGGAACACTGCTTGTCACTCAAAATTTAAACTTAAACCAAGGAAATCTCTAATTATTAATATGATTACCCCTTATTCACTGGCATTAAACGGGATATATTAAAGGAATCAGGCTTAAATCTTGTGCTACTGCTAAAAGGGTGAGAGAAGATTAAATAGCAGTTGATTGGGGTTGAAAAGGAATTAAATGGATTTCGAGAGAAAATGAGATCTCGATTGCAAAAATTATTGAATATGGATTAAATAAATATTCTAAATTCCCTTAAATAGTAAAAGAAAAGTCTATAATGGTACAGGGATACAGTTGCTCTTTCTAAAGAGCCTTTTAAATATTAACTAAAGATGAATTGATAAAATTAAATATAAGTTACTGTTGATAAAATTGGTTGCTCCACTTAGGGGATATGCCTGTTGGGAGGGCTATGATTTTTAGAAAATGAAAAAAATAGGGGATGTTGCTTGTAAAGTGGCAAGGTCTTATAAGAGTTTATAATCTTTCAAGAGAGTATTAA

5290.55

GTAAATTCTAGCTGAGATTAATAAATTATCAGAGTATGCAAAGTCATAAGTTTAATAGATTGATACTAAGCGAA
TACCTTATTCTAAGAAGCCCAATTGGTTTTCCTCCAGCATCTTACAAAGATTGTTACAATTATACAGCTT
AATTGAAGCTGAAAAGCGAAATATAAAATTAAAAGCATAGTCCTATTAGCGATTCTGCTCATATTATAATGCA
CCCCCCCATTCTCTTTGATGTTTAAAGGTCAAATTGTTAATTGAAAGAGATTAAAAGGACTTCAG
TTTCTTCGGGTAATGATTCTCTATTGCAATTGCTGAGTTGTTAGGCAATTAAATAGCTTGTAAATTAAAT
GAATATTATGTTTAAATTAGGGCTTTAATATGCATTGTTGAAACCTCTGGATATAGCAGCGAGAATAAG
ATTACAGCACTAGATATGGCTTTTGTGAAATCTTATATAGAAAAGTTAAATTATGCTTAATATTCAATTCTT
TAAGTATTATTTATCCAAAGAGTAGAAATTAGGAAC TGCTTGTCACTAAAATTTTAAACTTAAACAAAG
AAATGCTTATTATTAATATGATTACCCCTTATTGAGATGGCATTAAAACGGGATATATTAGGAATCAGGCTTA
AACTTGTTGCTACTGCTAAAAGGGTGAGAGAAGATTATAGCAGTTGTTAGGGGTTGAAAAGGAATTAAATG
GATTGGAGAGAAAATGAGATCTCGATTGCAAAATTTATGAAATATGGATTAAATAATATTCTAAATTCC
TTTAATAGTAAAATTAAAAGAAAAAGTCTATAATGGTACAGTGGATACAGTTGCTTTCTAAAGAGCCTTT
TATTATATTAACTAAAGATGAATTGATAAAATTAAATATAAGTTACTGTTGATAAAATGGTTGCTCCACTTA
GTGGGGATATGCCGTTGGGAGGGCTATGATTTTTAGAAAATGAAAAATAGGGGATGGCTTGTGTTAGTGG
CAAGGTAAGAGATTAGGGTTGGCAAGGTCTTATAAGAGTTTATAATCTTTCAAGAGAGTATTAA

E291.aa

MNSYDFITALVPIILIIIGLGIKKPAYYVIPISLIATVAIVIFYKNLGVNTSLAMLEGALMGIWIATVIIAAI
FTYKMSEDQKDIETIKNLSNVSSDRRIIVLLVAWFGNFLEGVAGYGTAVAIPVSILIAMGFEPFFACLICLIMN
TSSTAVGSGVIPITSLAQATNLDVNIVSSEIAFQLILPTLTIPFVLVILTTGGGIKGLGVFLTLLSGMSMAISQV
FISKTLGPELPAILGSILSMTITIVYARFFGNKETTERQSKNTISLSKGIACSPYILIVTFIVLVSPLFNKIHEY
LKTTFQSTISIYPEANPLHFKWIISPGFLIILATTISYSIRGVPMLKQLKIFTLTLKKMALSSFIICIVAIISRLMT
HSGMIPRDLANGISIITGKFGPLFSPLIGAIGTFLTGSDTVSNVLFGPLQTQMAENIGANPYWLAAANTTGATGGKM
ISPONITIATTTAGLIGQEGKLLSKTIIYALYYILATGLLVYL

.t291.aa.

QKDIEITKNILSNVSSDRRIVLLVAWGFNFLEGVAGYGTAVAI PVSILIAMGFEPFFACLICLIMNTSSTAYGS
VGIPITSLAQATNLDVNIVSSEIAFQLILPTLTI PFLVIL TGGGIKGLKGVLFLTLLSGMSMAISQVFISKTLP
ELPAILGSILSMTITIVYARFFGNKETTERQSKNTISLSKGIIACSPYI LIVTFIVLVSPLFNKIHEYLKTQSTI
SIYPEANPLHFKWII SPGFLIILATTISYSIRGVPMKLQLKIFTLKKMALSSIIICIVAI SRLMTHSGMIRDL
ANGISIITCKFGPLFSPLIGAITGFTLTGSDTVSNVLFGPLQTQMAENIGANPYWLAAANTTGTGGKMIS PQNITI
ATTTAGLICQEGKLLSKTIIYALYYILATGLLVYLV

291 nt

ATGAATTCTTATGATTTATAACAGCTTGGTACCAATAATCCTAATAATTATTGGACTTGGCATAATAAAGC
CAGCTTACTATGTAATACCCATATCATTAATAGCCACCGTGTCTAGTTATTTATAAAACTTGGGAATAGT
AAACCAAGTCTTCAATGCTTGAGGCGCTTAATGGGGATATGGCCAATAGCACTGTAATTATTGCTGCCATA

TABLE 1. Nucleotide and Amino Acid Sequences

TTTACATACAAAATGTCAGAAGATCAAAAAGATATAGAAAATTTAAACGTATCTTCTGATA
 GAAGAATTATAGTATTACTAGTGCATGGGGATTGAAATTTCAGAGGAGTTGCTGGATATGAACTGCTGT
 TGCAATTCCGTATCAATAATTAGCAATGGGATTGAAACATTTCGCTGCTTAACTGTTAATAATGAAC
 ACCTCATCAACCGCCTACGGATCTGGGATCCCTATAACATCTTAGCTCAAGCAACTAACTGGATGTTAAC
 TTGTTTCATCTGAGATTGCACTTCAACTAACTTCAACCTTAACAATACCTTTGACTGGTAATTCTTACAGG
 AGGGGGCATAAAGGATTAAGGAGTATTCTTACCTTAACAGGATGTCAATGGCAATATCTCAAGTA
 TTATATCAAAAACCTTGGTCCAGAACTTCCGCAATCCTTGAAGCATTCTTCTATGACAATAACAATAGTT
 ATGCAAGGTTTTGAAATAAGAAACTACTGAGGCCAAGCAAAACACAATATCCTTATCAAAAGGAAATTAT
 TGCCGTGCTCACCCCTACATTTAATAGTAACCTTATAGTGTGTTATCTCTCTTTAACAAAATTGATGAAATAC
 CTAACAACTTTCAAGCACTATTAGCATTATCCAGAACGAAATCCCTACACTTAAATGGATTATCTCCGG
 GCTTCTGATTATACTGCAACAAACATATCCTATTCAATACGGGGAGTCCAATGTTAAACAGCTAAAATATT
 TACATTAACCTGAAAAAAATGGCATTATCTCCTTATAATCATATGATTGCAATATCAAGATTAATGACA
 CATTAGTGGAAATGATAAGAGATCTGCTAATGGAATCTCAATAATAACAGGAAATTGGACCATTATTTAGCCCAC
 TAATTGGAGCTATTGGACATTTAACAGGAAGTGTACAGGTTCAAAATGTTCTTTGGACCTTACAAACACA
 AATGGCAGAAAATATTGGAGCAAATCCTTACTGGCTTGAGCAGCAAATACAACAGGAGCAACTGGAGGAAAATG
 ATTCTCCCCAAAACATCACAAATAGCAACAAACTGCTGGATTAAATTGGACAAGAAGGCAAGCTTTATCAAAAAA
 CAATAATTATGCTTATACTACATTAGCAACAGGATTGCTAGTTATTAGTATAA

t291.nt

CAAAAAGATATAGAAAATTTAAACGTATCTTCTGATAGAAGAATTATAGTATTACTAGTTG
 CATGGGGATTGAAATTTCAGAGGAGTTGCTGGATATGAAACTGCTGTTGCAATTCTGTATCAATATTAAT
 AGCAATGGGATTGAAACCATTTCGCTGCTTAACTGTTAATAATGAAACACCTCATCAACCGCCTACGGATCT
 GTGGGAATCCCTATAACATCTTAGCTCAAGCAACTAACTGGATGTTAACATTGTTATCTGAGATTGATTCC
 AACTAATACCTCAACCTTAACAATACCTTTGACTGGTAATTCTACAGGGAGGGGGCATTAAAGGATTAAAAGG
 AGTATTCCCTTACCTTACTCTCAGGAATGTCATGGCAATATCTCAAGTATTATATCAAAACTTGGTCCA
 GAACCTCCGCAATCCTGGAAAGCATTCTTCTATGACAATAACAATAGTTATGCAAGGTTTTGGAAATAAAG
 AACTAATGAGGCCAAGCAAAACACAATATCCTTATCAAAAGGATTATGCTGCTCACCCATCATTAAAT
 AGTAACCTTATAGTGTGTTATCTCTCTTTAACAAAATTGATGAAATACCTAAACACTTTCAAAGCACTATT
 AGCATTATCCAGAACAAATCCCTACACTTAAATGGATTATCTCTCCGGCTTCTGATTATACTTGAACAA
 CAATATCCTATTCAATACGGGAGTCCAATGTTAAACAGCTAAAATATTACATTAACCTTGAACAAAATGGC
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 GCTAATGGAATCTCAATAAACAGGAAATTGGACCATTATTTAGCCCCTAAATTGGAGCTATTGGACATT
 TAACAGGAAGTGTACGGTTCAAATGTTCTTTGGACCTTACAAACACAATGGCAGAAAATATTGGAGCAA
 TCTTACTGGCTTGAGCAGCAAATACAACAGGAGCAACTGGAGGGAAAATGATTCTCCCCAAAACATCACAAATAG
 CAACAACAATGCTGGATTAAATTGGACAAG

f296.aa

MPSPIRVFFLVLIFIFNPVLIAMLFILEPFLILFSFLGVFRIYFTRDYSRSREFEFYKLSFLLMAKLLSIL
 GTVTGEQLNYVNFIINSLNLSERGKSELYTIFHSAITKNNNADKILYTLKLGYFQHKDLFIWLFATLKEINRLSRY
 KNLEAEKFISYVGVFLLESDGYEAYKDINIKIVNPYSLGLTYSASDDEVKKAYKSLVIKYHPDKFANDPVRQKD
 ANDKFIKIQDAYEKICKERNIR

t296.aa

IYFTRDYSRSREFEFYKLSFLLMAKLLSILGTVTGEQLNYVNFIINSLNLSERGKSELYTIFHSAITKNNNADK
 ILYTLKLGYFQHKDLFIWLFATLKEINRLSRYKNLEAEKFISYVGVFLLESDGYEAYKDINIKIVNPYSLGLTYS
 SASDDEVKKAYKSLVIKYHPDKFANDPVRQKDANDKFIKIQDAYEKICKERNIR

f296.nt

ATGCCAAGCCCAATTAGAGTGTGTTAGTGTGTTGTTAATTTAATCCGTTAATAGCAATGC
 TTTTATTTATTCCTTATTTGATATTATTTAGTTAGGTGTTAGAATATACTTACAAGGGATTA
 CTCATATTCTAGATCTAGAGAGTGTGAATTAAACTTTCTTTTATTAATGGCTAAATTGCTATCTATTAA

TABLE 1. Nucleotide and Amino Acid Sequences

GGAACTGTAACGGGGAGCAGCTAAATTATGTCATTTATTATCAATTCTTGAATTGTCAGCTGGTAAAT
 CAGAATTGATACCATTTCTGCTATTACTAAAAATAATGCTGATAAAATTATACCCCTAAGCT
 TGGTTATTTCAAGCACAAGATCTTTATGGCTTTGCCACTCTAAAGAAATTAAACAGGCTTCTAGGTAT
 AAAAATTAGAAGCTGAAAATTATTCCTATGTTGGTGTGTTAGAACCTGAATCTGATGGTTATGAAGCTT
 ATAAAGATATTAATTTAAAGCTTAAACATATAGTGTGTTGGGTTAACATATAGTGTAGCGATGAGGT
 TAAAAGGCGTATAAAAGCCTGTTATAAAATATCCTGATAAGTTGCAAATGATCCTGTAAGACAAAAGAT
 GCAAATGATAAATTATAAAATTCAAGATGCTTATGAAAAAATTGCAAGGAAAGAAATATAAGGTAA

t296.nt

ATATACTTACAAGGGATTACTCATATTCTAGATCTAGAGAGTTGAATTTATAAACTTTCTTTTATTAATGG
 CTAATTGCTATCTATTTAGGAACGTAACTGGGGAGCAGCTAAATTATGTCATTTATTATCAATTCTTGA
 TTTGTCTGAACGTGGTAAATCAGAATTGATACCATTTCTGCTATTACTAAAAATAATGCTGATAAA
 ATTTTATATACCCCTAACGTTGGTATTTCAAGCACAAGATCTTTATGGCTTTGCCACTCTAAAGAAA
 TTAACAGGCTTCTAGGTATAAAATTAGAACGCTGAAAATTATTCCTATGTTGGTGTGTTAGAACCTG
 ATCTGATGGTTATGAAGCTTAAAGATATTAATTTAAATTGAAATCCTTATAGTGTGTTGGGTTAACATAT
 AGTGTAGCGATGATGAGGTTAAAGGCGTATAAAAGCCTGTTATAAAATATCCTGATAAGTTGCAAATG
 ATCCTGTAAGACAAAAGATGCAAATGATAAATTATAAAATTCAAGATGCTTATGAAAAAATTGCAAGGAAAG
 AAATATAAGGTAA

f3.aa

MKKKNLSIYIMILISLLSCNTSDPNELTRKKMQDKNVKILGFLEKIQADNKEIVEKIEKKEKQMVQAASVAPINV
 ESNFPYIQLQEEIEIKEELVPNTDEEKKAEKASDGSLFALKVDDENKLKNESAQLESSFNNVYKEILELADLIQ
 AEVHVAGRINSYIKKRKTTKEKEYKKREIKNIEKQALIKLFNQLLEKRGDIENLHTQLNSGLSERASAKYFFEKA
 KETLKAAITERLNNKRKNRPWWARRTHSNLAIQAKNEAEDALNQLSTSSFRILEAMKIKEDVKQLLEEVKSFLDSS
 KSKIFSSGDRLYDFLETSK

t3.aa

NELTRKKMQDKNVKILGFLEKIQADNKEIVEKIEKKEKQMVQAASVAPINVESNF PYIQLQEEIEIKEELVPNTD
 EEKKAEKASDGSLFALKVDDENKLKNESAQLESSFNNVYKEILELADLIQAEVHVAGRINSYIKKRKTTKEKEY
 KKREIKNIEKQALIKLFNQLLEKRGDIENLHTQLNSGLSERASAKYFFEKAETLKAAITERLNNKRKNRPWWAR
 RTHSNLAIQAKNEAEDALNQLSTSSFRILEAMKIKEDVKQLLEEVKSFLDSSKSKIFSSGDRLYDFLETSK

f3.nt

ATGAAAAAAAAATTATCAATTACATGATAATGCTAATAAGTTATTATCATGTAATACAAGTGACCCCAATG
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 AAATGAATCTGCGAATTAGAATCTAGTTAATAATGTTATAAGAAATCTTAGAAGCTGAGATTAAACAA
 GCAGAGGTGCATGGCAGGAAGGATAAATAGCTATAAAAGAAAAGACCACTAAAGAAAAGAATATAAGA
 AGAGAGAAATTAGAATAAGATAGAAAACAGGCTCTAATTAGTTGTCATCAGTTATTAGAAAAAGAGGCGA
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 AAAGAAAATTAAAGCTGCTATTACTGAAAGATTAATAACAAACGTAAAATCGGCCATGGTGGCAAGAAGAA
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 GATACTGAAAGCAATGAAAATAAGGAAGATGAAACAGCTTCTGAAAGAAGTAAATCTTTCTAGATTCTTCA
 AAGAGAAAATTCTTCTAGTGGCGATAGATTATGATTAGAGACGAGTAAATAA

t3.nt

• AATGAATTAACCTCGTAAAAAAATGCAAGACAAGAACGTGAAAATTAGGATTAGAGAAAATTCAAGCAGATA
 ATAAAGAAAATTGTTGAAAACATATAGAAAAAAAGAAAACAAATGGTGCAGGCTGCTTGTAGCACCTATTAA
 TGTAGAGAGTAATTCCCATTATCTCAAGAAGAAATAGAGATAAAAGAAGAGAGTTGGTCCAAATACTGAT

TABLE 1. Nucleotide and Amino Acid Sequences

GAAGAAAAGAAGGCAGAGAAGGCATAAGCGATGGGAGTCTTGAATTGCTAAATTAGTTGATGATGAAAATAAAC
 TAAAAAATGAATCTGCGCAATTAGAATCTAGTTAATAATGTTATAAAGAAATCTTAGAACTGCAGATTTAAT
 ACAAGCAGAGGTGCATGTTGCAGGAAGGATAAATAGCTATAAAGAAAAGACCACTAAAGAAAAAGAAATAT
 AAGAAGAGAGAAATTAGAATAAGATAGAAAAACAGGCCTAATTAGTTGTCATCAGTTAGAAGGAGAA
 GCGATATTGAAAATCTTCATACTCAATTAAATAGTGGACTTAGCGAGAGGACATCTGAAAATACTTTTGAGAA
 AGCCAAAGAAACTTAAAGCTGCTATTACTGAAAGATTAATAACAAACGTAAAATCGGCCATGGTGGCAAGA
 AGAACACATAGTAATTAGCAATACAGGAAAAATGAGGCAGAGGATGTTAAACCAATTAGTACTCTTCTT
 TTAGGATACTTGAAGCAATGAAAATAAGGAAGATGTTAAACAGCTTCTGAAGAAGTAAAATCTTTCTAGATTC
 TTCAAAGAGCAAAATCTTTCTAGTGGCGATAGATTATGATTTTAGAGACGAGTAAATAA

f30.aa

MNKKILTLVLILSISSVMLSKSITKKSKYKIIIRDYFINSNYVLVKIENKDLKFTISKPIYDKKLNNYFFKGQTT
 SHFLISNNVDIAINTSPYEVKQNMFFPKGLYIYNKKMISKQINNYGEIVIKHNKIIILNPKEDEIENCDYGFSGFFF
 LIKNGKYKKNFKETRHPRTIIGTDKNNKHLFLVTIEGRGVNNSKGASLNEAIDFALSYGMTNAINLDGGGSSTLVV
 KSNNAPYKLNFTANIFGQERPVFPFLGIKLPN

t30.aa

LSKSITKKSKYKIIIRDYFINSNYVLVKIENKDLKFTISKPIYDKKLNNYFFKGQTTSHFLISNNVDIAINTSPYEV
 KQNMFFPKGLYIYNKKMISKQINNYGEIVIKHNKIIILNPKEDEIENCDYGFSGFFFVLIKNGKYKKNFKETRHPRTI
 IGTDKNNKHLFLVTIEGRGVNNSKGASLNEAIDFALSYGMTNAINLDGGGSSTLVVKSNNAPYKLNFTANIFGQER
 PVFPFLGIKLPN

f30.nt

ATGAATAAAAAATATTAACACTGCTAGTATTGATTTAAGTATTCATCAGTACTAATGCTGTCAAATCAATCA
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 AGCCATTCTTAATTCTAACATGTTGACATTGCAATTAAACACAAGTCATACGAAGTTAAACAAAACATGTTT
 TCCCAAAAGGACTATACATATATAATAAAAAATGATTCAAAACAAATAAAACTACGGAGAGATTGTAATAAA
 GCACAAACAAAATTATTAATCCAAAGGAAGACGAAATAGAAAATGCGATTATGGATTAGCGGATTTTGT
 TTAATCAAAACGGAAAGTATAAAAAAATTAAAGAAACAAGGCACCCAAAGAACATAATAGGAACGTATAAAA
 ATAACAAGCATTATTTCTGTTAACATAGAAGGAAGGGGTGCAATAATAGCAAAGGGGCCTCTTAATGAAGC
 TATTGATTTCGATTAAGCTACGGCATGACTAACGCTATTAAATCTAGACGGGGGGGCTCAAGCACTCTTGTGTA
 AAATCAAATAACGCTCCTAACAAATTAACTTCACAGCAAACATCTTGGACAGGAAAGACCTGTCCCATT
 TAGGAATAAAACTTCCTAACATTGA

t30.nt

CTGTCCAAATCAATCACCAAAAAATCCAAATACAAAATTATTAGGGATTATTCATAAACAGCAATTATGTTCTGG
 TGAAAATTGAAAATAAGATCTAAATTCACCATATCAAAACCTATTTACGACAAAAGCTAAATAATTACTCTT
 TAAAGGCCAAACAACAAGCCATTCTTAATTCTAACATGTTGACATTGCAATTAAACACAAGTCATACGAAGTT
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 ATAGGAACGTATAAAATAACAAGCATTATTTCTGTTAACATAGAAGGAAGGGGTGTCATAATAGCAAAGGG
 CCTCTCTTAATGAAGCTATTGATTTCGATTAAGCTACGGCATGACTAACGCTATTAAATCTAGACGGGGGGGCTC
 AAGCACTCTGTTGAAATCAAATAACGCTCCTAACAAATTAAACTTCACAGCAAACATCTTGGACAGGAAAGA
 CCTGTCCCATTCTAACATTGAAATAAAACTTCCTAACATTGA

f308.aa

MQLLKNKYPFKRALLDLFLVYAIYVLASPFVNVNSEFWNVDENHFYFWISRSFLIIIFIYFFKLTSSYDDFRVEFF
 IPKFKFIFLWDSVLIFIKTILIAMIVIFLLEYLLPESVLVYYFQNNAGFNWKISSKKAFFLMTFTSFFTGA

TABLE 1. Nucleotide and Amino Acid Sequences

EELFYRAFVITKFTQMGFPVVATAILSSMFFAYGHLYYGINLGLVTFILGIGFAFTYLRYKNVYYVIFIHSFYNIIVSSLLLFLN

t308.aa

NSEFWNVDENHFYFWISRSFLIIFIIYFFKLTSSYDDFRVEFFIPKFKFIFLWDSVLIFIKTILIAMIVIFLIAFL
LEYLLPESVLVYYFQNNAGFNWKISSKKAFFLMTFTSFFTGAEEELFYRAFVITKFTQMGFPVVATAILSSMFFAY
GHLYYGINLGLVTFILGIGFAFTYLRYKNVYYVIFIHSFYNIIVSSLLLFLN

f308.nt

ATGCAATTGTTAAAAAATAATCCATTCAAGCGGGCTTGCTGATCTTTGGTCTATGCTATTGTTATT
TGGCATCTCCTTTGTAATGTTAATTCAAATTTGGAATGTTGATGAAAATCATTTATTGGATTCAG
ATCTTTTAATTATTTATAATTAACTTACCAAGTTCTATGATGATTAGAGTAGAGTTTT
ATTCCATAATTAAATTATTTCTTGGGATTCTGTTAAATTAAACAAATTGATTGCAATGATAG
TCATTGTTAATAGCTTTGCTGAATATTGTTGCCAGAATCGGTACTTGTCTATTATTCAAAACAATGC
TGGATTTAATTGGAAGATTAGCAGTAAAAAGCATTGTTAATGACTTTACCTCTTTTACAGGAGCTTT
GAAGAACTTTTACAGGGCTTTGTTACTAAGTTACACAAATGGGATTCTGTTAGCTACCGCCATT
TTAGTAGTATGTTTTGCTATGGCATTATATTATGGAATTAGGATTGTTACATTATATTAGGGAT
ATTGGTCTTTACTTAAAGGTATAAAATGTATATTATGATTTACATAGTTTATAATATTATT
GTTAGCAGCTTGTGCTTTGTTGAATTAA

t308.nt

AATTCAAATTTGGAATGTTGATGAAAATCATTGTTATTGGATTCAGATCTTTAATTATTGTTATAA
TTTATTTTAAACTTACCAAGTTCTATGATGATTAGAGTTAGCTACCTCTTAAATTAAATTATT
TCTTGGGATTCTGTTAATTAAACAAATTGATTGCAATGATAGCTATTGTTAATAGCTTTTG
CTGAATATTGTTGCCAGAATCGGTACTTGTCTATTGTTAAATGGAAGATTAGCA
GTAAAAAAGCATTGTTAATGACTTTACCTCTTTACAGGAGCTTGAGAAACTTTTACAGGGCTTT
TGTTATTACTAAGTTACACAAATGGGATTCTGTTAGCTACCGCCATTCTAGTAGTATGTTTGCTTAT
GGGATTATATTATGGAATTAGGATTGTTACATTATATTAGGGATATTGTTACTTATTAA
GGTATAAAATGTATATTATGATTTACATAGTTTATAATATTGTTAGCAGCTGTTGCTTTTT
GAATTAA

f31.aa

MKKYLFFILFLISSNNLIVSYPLSFGGGSYQFTNYTDKTGATKFAPNFTRADHGINLNLFDDANYVLFEMSYKEA
FVVTNGRYFSLGLYGTYPMVKEQVRMLFPLIGFKYAFDLSSNNFLFLSMGLAADLFIPDLDGLYIRPLFMLS
ISPFSNYKNFSGLTTEIMLGFNIGWRFFN

t31.aa

IVSYPLSFGGGSYQFTNYTDKTGATKFAPNFTRADHGINLNLFDDANYVLFEMSYKEAFVVTNGRYFSLGLYGT
YPMVFKEQVRMLFPLIGFKYAFDLSSNNFLFLSMGLAADLFIPDLDGLYIRPLFMLSISPFSNYKNFSGLTTEI
MLGFNIGWRFFN

f31.nt

ATGAAGAAATATCTTTTTATTTATTCTCATCTCTTAATAATTAAATTGTTCTTATCCACTTCTTTG
GTGGAGGTTTTCTTATCAATTACTAATTACTGATAAAACAGGCCTTCAAAATTGCTCAAATTACAG
AGCAGATCATGGGATAATTGAATTATTTGATGCAAATTATGACTTTGAAATGCTTACAAAGAGGCT
TTTGTGTTACTCACAATGGGAGATAATTCTCGCTGGCTTATGGAACATATCCAATGGTTCAAAAGAGCAGG
TTAGAATGCTTCCATTAAATGGCTTAAATATGCTTTGATTTAAGCTCTAAACTCAATCTCTTTTT
AAGCATGGGCTGCTGATCTTTATTCCGATCTGATGGTTATATATTAGGCCTTGTATGCTTCT
ATTCTCCATTCTAATTATAAAATTCTGGGTTAACACTGAGATTGCTGGATTAAATCGGTTGA
GATTTTCAATTAG

TABLE 1. Nucleotide and Amino Acid Sequences

t31.nt

ATTGTTCTTATCCACTTCTTGGTGGAGGTTTTCTTATCAATTACTAATTATACTGATTAACAGGGCCAA
 CTAATTTGCTCAAATTTACAGAGCAGATCATGGGATTAATTGAATTATTTTGTCAATTATGACT
 TTTGAAATGTCTTACAAAGAGGTTTGTTACTCACAATGGGAGATAATTCTCGCTGGGCTTATGGAA
 TATCCAATGGTTTCAAAGAGCAGGTTAGAATGCTTCCCATTAAATTGGGTTTAATATGCTTGTCAATT
 CTAATAACTCAATCTCTTTTAAGCATGGGCTGCTGATCTTATTCCGATCTGATGGTTAA
 TATTAGGCCTTGTATGCTTCTATTCTCAATTATAAAATTCTGGGTTAACACTGAGATT
 ATGCTTGGATTTAATATCGGTTGGAGATTTCAATTAG

f939.aa

MKQKYENYFKKRLILNLLIFLLLACSSESISQLGNLQKIKHEYNILGSSSPRGISLVGETLYIAAMHLFKKENGK
 IEKIDLSNSYEFINDIVNISGKTYLLAQNKEEELEVCELNKGDWTLKFKPLKAYKFLKSVGRDGVEAYILAIDK
 NRREKIFDLOQSDKTPPQATENDKFYQISNEENLITGNSLKIWQMNNTYTNIDYQQAKEIMPIIKTSIRGSSEV
 VMTGGYNNLDTKFKVYSNTNNYTPPIFPIQDEVGEFSSYFAREFNDAILIGSNNGFAEFTKNEGIFALRAPS
 PGAYNGSQLSKTGLNDIIPVSNNTIYILTQGKGLWKLNRKLTKE

f939.aa

CSSESISQLGNLQKIKHEYNILGSSSPRGISLVGETLYIAAMHLFKKENGKIEKIDLSNSYEFINDIVNISGKTY
 LLAQNKEEELEVCELNKGDWTLKFKPLKAYKFLKSVGRDGVEAYILAIDKNNREKIFDLOQSDKTPPQATENDK
 FYQISNEENLITGNSLKIWQMNNTYTNIDYQQAKEIMPIIKTSIRGSSEVLMTGGYNNLDTKFKVYSNTNNY
 PPIQDEVGEFSSYFAREFNDAILIGSNNGFAEFTKNEGIFALRAPS
 KVEPGAYNGSQLSKTGLNDIIPVSNN
 IYILTQGKGLWKLNR
 KLTKE

f939.nt

ATGAAACAAAATACGAAAATTTAAAAAGATTAATTAAACCTATTAAATTACTACTAGCACTGCT
 CAAGCGAATCCATATTTACAATTAGGAAATCTGCAAAAATAAAACATGAATACATTATTTGGCAGTTCAAG
 TCCAAGAGGAATTCTCTAGTAGGAGAAACTCTCTACATTGCAGCCATGCATTATTAAAGAAAACGGCAAG
 ATTGAAAAATTGATTGAGCAATTCTTATGAGTTATAAACGACATTGTAAATTATCTGGAAAACCTATCTT
 TAGCGAAAACAAGAAGAATTAGAAGTTGGCAGCTAAATGGAAAAGATTGGACATTAAATTAAAC
 GCTAAAAGCATATAATTCTAAAATCCGTAGGAAGAGATGGCGTAAAAGAAGCATAATTAGCTATAGATAAA
 AATAATCGTGAGAAAATTGATCTACAAGGATCTGACAAAACACCACAAAGCTACTGAAAATGACAAATT
 ATCAAATATCAAATGAAGAAAATTACAGGAAATTCACTCAAATATGGCAAAATGAAATTACATACAGC
 AAACATAGACTATCACAGGCCAAGAAATAATGCCATCATAAAACAAGCATTAGGGCTCTCTGAGTT
 GTAATGACTGGTGGTTACAATAATTAGATACAAAATTAAAGTTACTCAATTACAAATTACACAGGCCAA
 TATTATTCAAGACGAAGTAGGCGAATTAGCAGCTACTTGCAGAGAAATTAAATGATGGCAATTAGCTGGAG
 TAATAATGGATTGAGAATTACAAAAATAAGAAGGAATTGGCCTACGGGCACCCCTAAAATCTGTAGAA
 CCTGGAGCTTATAACGGATCTCAGCTAACGAAAACAGGCCATTAGTATATTCTCTGATCAAAACA
 ACATATTAACTCAGGCCAAGGGTTGTGAAATTGAAAACAGAAAATTACTAAAGAATTAA

t939.nt

TGCTCAAGCGAATCCATATTTACAATTAGGAAATCTGCAAAAATAAAACATGAATACATTATTTGGCAGTT
 CAAGTCCAAGAGGAATTCTCTAGTAGGAGAAACTCTCTACATTGCAGCCATGCATTATTAAAGAAAACGG
 CAAGATTGAAAAATTGATTGAGCAATTCTTATGAGTTATAAACGACATTGAAATTATCTGGCAAAACCTAT
 CTTTAGCGCAAAACAAGAAGAATTAGAAGTTGGCAGCTAAATGGAAAAGATTGGACATTAAATT
 AACCGCTAAAAGCATATAATTCTAAAATCCGTAGGAAGAGATGGCGTAAGAAGCATTAAATT
 TAAAATAATCGTGAGAAAATTGATCTACAAGGATCTGACAAAACACCACCAAGCTACTGAAATTGACAA
 TTTTATCAAATATCAAATGAAGAAAATTCAATTACAGGAAATTCACTCAAATTGCAAAATGAAATTACAT

TABLE 1. Nucleotide and Amino Acid Sequences

ACACAAACATAGACTATCAACAGGCCAAGAAAATAATGCCTATCATTAAAACAAGCATTAGGGCTCTCTGAAGT
 TTTAGTAATGACTGGGTTACAATAATTAGATACAAAATTAAAGTTACTCAAATACAAATAATTACACAACG
 CCAATATTATTCAAGACGAAGTAGGCGAATTAGCAGCTACTTGCAAGAGAATTAAATGATGCGATATTAAATCG
 GAAGTAATAATGGATTGCAGAATTACAAAAATAAAGAAGGAATTTCGCCCTACGGGCACCCCTAAACATCTGT
 AGAACCTGGAGCTTATAACGGATCTCAGCTAAGCAAAACAGGCTTAATGATATTATTCCGTATCAAACACAG
 ATTACATATTAACTAGGGCAAGGTTGTGGAAATTGGAAAACAGAAAATTAACTAAAGAATAA

f739.aa

MQSGLKIKLILFFCCFACSCDINYPEIKELDYKINYYFTENRLDYSMSFDFAIKVINSKDVFKLSIENKNTNEFIQ
 VVINNNYSSFFIDSSLGKDILYCKDLRFNFFDKTFEDFTSCVRLFDKGMRVYNRELVISLGMSKYDLDDVHNVYVYKS
 KDMEMLNKLSNSKVFFVKTYPDKLHPVSSVVRIDSIDILEIDKAFDNYISFYVVEKNSNLFFKVG

t739.aa

CCFACSCDINYPEIKELDYKINYYFTENRLDYSMSFDFAIKVINSKDVFKLSIENKNTNEFIQVINNNYSSFFIDS
 SLGKDILYCKDLRFNFFDKTFEDFTSCVRLFDKGMRVYNRELVISLGMSKYDLDDVHNVYVYKS
 VFFVKTYPDKLHPVSSVVRIDSIDILEIDKAFDNYISFYVVEKNSNLFFKVG

f739.nt

ATGCAGAGCGGATTAATAATTAAATTATTGTTTTGTTGCTTGCACATAAATTATCCGG
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 TGCAATTAAAGTTATAAAATTCAAAAGATGTTTAAATTATCAATAGAGAATAAGAACACTAATGAGTTATTCAA
 GTGATTAATAATAATTATAGCTTTTTATTGATTCTAGCCTGGAAAGGATATTCTATATTGTAAGGATTGAA
 GGTTTAATTGATAAAACTTTGAAGATTTCACCTCATGTTGCTTTGATAAGGGCATGAGAGTATA
 CAATAGAGAGCTGTTATTCTTGGGTATGTCAAATATGATTAGATGTTCACATTATGTATATAAGTCT
 AAAGATATGAAATGTTAAACAAGTTAACGAAATTCCAAAGTATTGTTAAACTTATAAGACAAACTACATC
 CGGTCTCTCAGTTGTTAGAATTGATTCAATAGATATTCTAGAGATTGATAAGCATTGATAATTAGATT
 TTATTATGTCGAAAAAAATTCAAATCTTTTAAAGTTGGCTGA

t739.nt

TGTTGTTTGCTTGTCTTGCACATAAATTATCCGGAGATAAAAGAGCTGATTATAAGATAAAATTATTATTAA
 CTGAAATCGCTTAGATTACTCTATGAGTTTGATTTGCAATTAAAGTTAAATTCAAAAGATGTTTAAATT
 ATCAATAGAGAATAAGAACACTAATGAGTTATTCAAGTGATTAAATAATAATTATAGCTTTTTATTGATTCT
 AGCCTGGAAAGGATATTCTATATTGTAAGGATTGAGGTTAATTGATAAAACTTTGAAGATTACCT
 CATGTTGCTCTTTGATAAGGGCATGAGAGTACAATAGAGAGCTGTTATTCTTGGGTATGTCAAATA
 TGATTAGATGATGTCACAATTATGTATATAAGTCTAAAGATATGAAATGTTAAACAAGTTAACGAAATTCCAA
 GTATTGTTAAACTTATAAGACAAACTACATCCGGCTCTTCAGTTGTTAGAATTGATTCAATAGATATT
 TAGAGATTGATAAAGCATTGATAATTACATAAGTTTATTATGTCGAAAAAAATTCAAATCTTTTTAAAGT
 TGGCTGA

f742.aa

MNKKHTNFSVLLLLIFLLILSFGGFGYYIYQSKLNDKNREIMLNEVKNSVIDRNYKKAYSVAKLLQDKYPQNEDIA
 MLNTNLAELIANSSPFESKDLQRDSANQILDKIKGQDNTKTNVNFIAFNNRYIKDSTITENYSDRNDVGIEDE
 DISEFKKSKEPEKIKPNTNPKEEDQIIQSPNPKLSVNDQKNLENLEKLKKNLSGKSNSENILNDSQKIENDKQNTN
 LSKEKNSENILKTPDNSKYSNNNNNTSLKKISSNSQKESELSPPSQTIIGKIQYRPSYLIKELYEILDDINTGRV
 TLGKNRLKELIKKGLSNKFQKVNELIENSKNKEASNLLTLIKKDEPNLINIPKDPYKEIFQLDKEDKKPQYLE
 DLKSKVHSIKPIDLENTKSRQQAIKDLNEFLKNNPNDQAQASKTLAQANKIQHLEDLKSKVHSIKPIDLENTKSRQQ
 AIKDLNEFLKNNPNDQAQASKTLAQANKIQHLEDLKSKVHSIKPIDLENTKSRQQAIKDLNEFLKNNPNDQAQASKT
 LAQANKIQHLEDLKSKVHSIKPIDLENTKSRQQAIKDLNEFLKNNPNDQAQASKTLAQANKIQHLEDLKSKVHSIKPI
 DLENTKSRQQAIKDLNEFLKNNPNDQAQASKTLAQANKIQHLEDLKSKVHSIKPIDLENTKSRQQAIKDLNEFLKNN
 PNDQAQASKTLAQANKIQHLEDLKSKVHSIKPIDLENTKSRQQAIKDLNEFLKNNPNDQAQASKTLAQAYENNGDLLK
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TABLE 1. Nucleotide and Amino Acid Sequences

QIDKNYGTAYYQKGIAECKNGDMQQAFASFKNAYNLDKNPNYALKAGIIVSNLGNFKQSEELYNFFNANAKKPNEIAIYNLSIAKFENNLEESLETINKAIDLNPBKSEYLYLKASINLKKENYQNAISLYSLVIEKNPENTSAVINLAKAYEKSGNKSQAISTLEKIINKNNKLALNNLGILYKKEKNYQKAIEIFEKAIINSDIEAKYNLATTIEINDNTRAKDLLREYTKLKPNNPEALHALGIIEYNENNNDOTLRELIKFPNYYKKNENIKKIIGI

t742.aa

KLNDKNREIMLNEVKNSVIDRNYKKAYSAKLLQDKYPQNEDIAMLTNTLAEIANSSPFESKDLQRDSANQILD
KGQD
NTKTNVNENFDIAFNNRYIKDSTITENYDRNDDVGIEDEDISEFKKS^KPIPEKIKPNTNPKEEQDQI^IQSPNPKLSV
NDQKNLNLEKLKKNLSGKNSENILNDSQKIEENDKQNTNL^SKEKNSENILKTPDNSKYSNNNNNTSLKKISSNSQ
KESELSPPSQ^TI^GKIYRPYSYLIKELYEILDINTGRVTLGK^NRL^KELIK^GLSNKFQKVNEI^ENSKNKEASN
LLLTLIKK^DIEP^NLN^IIPKDPYK^KE^IFQ^LD^KE^DKKPQYLEDLK^SKVHS^IK^PIDL^ENTKS^RQQAI^KDLNEFLKNNP
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KVHS^IK^PIDL^ENTKS^RQQAI^KDLNEFLKNNP^NDAQASKT^LAQANKI^QHLEDLK^SKVHS^IK^PIDL^ENTKS^RQQAI^KD
LNEFXKNNP^NDAQASKT^LAQANKI^QHLEDLK^SKVHS^IK^PIDL^ENTKS^RQQAI^KDLNEFXKNNP^NDAQASKT^LAQANKI^QHLEDLK^SKVHS^IK^PIDL^ENTKS^RQQAI^KD
TKS^RQQAI^KDLNEFXKNNP^NDAQASKT^LAQAYENNGD^LKAENAYEK^II^IKLNTQEDHYKLG^IIRFKLKKYEHSIE
SFDQT^IKLD^PKH^KKALHNKG^IALMMLNKNK^KAI^ESEFK^AI^IQIDKNYGT^AYYQKG^IAE^EKGMDMQAFAS^FKNAYNL
DKNP^NYALKAGIVSNNLGNFKQ^SEEYL^NFFNANAKPNEIAIYNLSIAKFENN^KLEESLETINKAIDLN^PEKSEYL
YLKASINLKKENYQNAISLYSLVIEKNPENT^SAYINLAKAYEKSGNK^SQA^ISTLEK^II^IINKNNKLALNNLG^ILYK^E
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IKKFP^NYKKNENIKK^IIGI

f742.nt

ATGAAATAAAAACATACAAATTTCGGTATTATTGCTTTAATTTCTTACTTATCTTACATTGGGGCTTG
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GCCCTCTAAACTTAGCTCAAGCTAATAAAATACAACACCTAGGACCTTAAATCTAAGGTTATTCAATAAAAC
CCATTGATCTGAAAACACAAAT
CACGCCAACAGCCATTAGGATCTAACGAAATTCTTAAACAAATCCCAATTGACGCCAGGCCCTCTAAACTTT
AGCTCAAGCTAATAAAATACAACACCTGGAGGACCTTAAATCTAAGGTTATTCAATAAAACCCATTGATCTGAA
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TABLE 1. Nucleotide and Amino Acid Sequences

TAAAACCTTAGCTAAGCTTATGAAAACAATGGAGATTGCTAAAAGCAGAAAATGCATACGAAAAAATTATCAAA
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 TACCAAAAAGGAATAGCAGAAGAAAAATGGCGATATGCAACAAGCATTGCAAGCTTAAAATGCCTACAATC
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 TGCAACCACCTCAATTGAAATTGATAACACAAGAGCTTAAAGACCTTCAAGAGAATATACAAAATTAAAACCA
 ACAATCCAGAGGCCATTACATGCACTAGGAATAATAGAATAATGAAAATAACATGATCAAACACTAAAGAGAAC
 TTATAAAAATTCCAATTACAAAAAAATGAAAATTAAAAATAATAGGAATATAA

t742.nt

AAATTAAATGACAAAATCGAGAAATAATGCTAAACGAAGTTAAAATAGCGTAATAGATCGAAACTATAAAAAG
 CATATTCTGTTGCAAAACTTCTGCAAGACAATAACCCAAAATGAAGACATTGCAATGCTTACAAATACACTAGC
 AGAAATTGCCAACAGTAGTCCTTTGAATCAAAGACTTGCAGAGATCTGCTAATCAAATCTTAGACAAGATC
 AAAGGTCAAGACAATAACAAAACAAATGTAACGAAAATTGATATAGCATTAAATAATAGATACATTAAAGACA
 GCACAATAACAGAAAACTACTGACAGAAACGATGATGTTGCCATTGAAGATGAAGACATATCTGAATTAAAAA
 AAGCAAAATCCCAGAAAAATAAAACCAATAACAAACCAAAAGAAGAAGACCAAATAATACAATCTCCAAATCCG
 AAATTAAAGTGTAAATGACCAAAAAATTATTAAATTGGAAAAGTAAATTTAAAGTGGAAAATCAAATA
 GTGAAAATATTAAACGATTCTCAAAAATAGAAAATGATAAGCAAAACCAAATTATCCAAAGAAAAATTTC
 GGAGAATATTAAACCTCGGACAACAGTAATATTCAAACAATAACAAACTACATCTTAAAAAAATTCT
 TCAAATTCCAAAAGAAAGTGTGAGCTTCTCCACCCAGTCACAAACAAATAATAGGAAAATTATAGGCCATATAGCT
 ACTTGATAAAAAAGAGCTATGAAATTAGACGATATTAAACCGGAAGAGTCACACTTGGAAAAACAGATT
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 TTCAATAAAACCCATTGATCTGAAAACACAAATACGCCAACAGCATTAGGATCTAAACGAATTCTTGAAA
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 GACCTTAAATCTAAGGTTATTCAATAAAACCCATTGATCTTGAACACAAATACGCCAACAGCATTAGG
 ATCTAAAGAATTCTTAAAAACAAATCCCAATTGACGCCAGGCCCTCTAAACTTAGCTCAAGCTAATAAAATAAC
 ACACCTGGAGGACCTTAAATCTAAGGTTATTCAATAAAACCCATTGATCTTGAACACAAATACGCCAACAA
 GCCATTAAAGGATCTAAACGAATTCTTAAACAAATCCCAATTGACGCCAGGCCCTCTAAACTTAGCTCAAGCTAAT
 AAAATAACACACCTGAGGACCTTAAATCTAAGGTTATTCAATAAAACCCATTGATCTTGAACACAAATCAG
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 AGCTAATAAAATAACACACCTAGAGGACCTTAAATCTAAGGTTATTCAATAAAACCCATTGATCTTGAACACAA
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 CTTAGCTCAAGCTAATAAAATAAC
 AACACCTGGAGGACCTTAAATCTAAGGTTATTCAATAAAACCCATTGATCTTGAACACAAATCACGCCAAC
 AGCCATTAAAGGATCTAAACGAATTCTTAAACAAATCCCAATTGACGCCAGGCCCTCTAAACTTAGCTCAAGCTT
 ATGAAAACAATGGAGATTGCTTAAAGCAGAAAATGCATACGAAAAATTATCAAACCTACAAATACCAAGAAGA
 TCACTATAAAACTTGGAACTTCAAGCTTAAAAGTATGAACACTCAATAGAATCATTGATCAAACAAATA
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 GCATTAAAGCAGGAATAGTATCAAATAACTTGGCAACTTCAACAAAGTGAAGAGTATTAAATTGTTAATG
 CCAATGCCAAAAACCTAACGAAATTGCTATTACACCTATCAATAGCAAAATTGAAAACAATAACTGAAGA
 ATCTCTTGAACAAATAACAAAGCCATTAGATTAAATCCAGAAAAAGTGAATATTATAATTAAAAGCATCTATA
 AATCTTAAAAAGAAAATTACCAAAATGCTATATCAGCTTACAGCTTAGTAATTGAAAAACCCGTAAAATACTT

TABLE 1. Nucleotide and Amino Acid Sequences

CAGCCTATATAAACCTGGAAAAGCATATGAAAAATCAGGAATAAAAGTCAGCAATCTCAACTCTTAAAAGAT
 AATAAACAAAATAATAATTAGCCTAACAAATCTTGGGATACTTTACAAAAAGAAAAAAATTATCAAAAAGCA
 ATTGAAATTTGAAAAGCAATAATCAATTAGATATTGAAGCAAAATAATCTGCAACCCTAATTGAAA
 TTAATGATAACACAAAGAGCTAACAGACCTCTAAGAGAATATAACAAAATTAAAACAAATCCAGAGGCCTTACA
 TGCACTAGAATAATAGAATATAATGAAAATAACATGATAACACTAAGAGAACTATAAAAAATTCCAAATT
 ACAAAAAAAATGAAAATATTAACAAATAATAGGAATATAA

f743.aa

MRIYLFLNKNYKIFILFLILILNSKLAWSQLRLIRIGKEEMKNKYIQAIETLSDAIKKYPKVQLGYYFLSIAYREN
 NQLTEAEGALLDGIAVGEIDYIYLGYELGNIMFNRGEYYPLAIKYYSNSIKSRPNYDSALLNRANAYVQQGKITS
 KEKEYQKAWDSYTMIAHDYSQFITLRSKTEKKDSILLIISYLRNEKINLEQLDKSLKGRTEHIVYAKEDKNQILKD
 SFKDNLETNSLIELEKLNWQEELYIDE

t743.aa

YSQRLIRIGKEEMKNKYIQAIETLSDAIKKYPKVQLGYYFLSIAYRENNQLEAEGALLDGIAVGEIDYIYLGYE
 LGNIMFNRGEYYPLAIKYYSNSIKSRPNYDSALLNRANAYVQQGKITSKEKEYQKAWDSYTMIAHDYSQFITLRS
 KTEKKDSILLIISYLRNEKINLEQLDKSLKGRTEHIVYAKEDKNQILKDSFKDNLETNSLIELEKLNWQEELYIDE

f743.nt

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 TGGC
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 AGTGTGCTATTAAAAATATCCAAAAGTACAACCTCGGCTATTACTTTTATCAATAGCATAACAGAGAAAATAATC
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 TTAAAGACAACCTAGAAACAAATTCTTAATTGAGCTAGAAAACCTTAATTGGCAAGAGGAGTTACATAGATGA
 ATAA

t743.nt

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 TAAAGACAACCTAGAAACAAATTCTTAATTGAGCTAGAAAACCTTAATTGGCAAGAGGAGTTACATAGATGAA
 TAA

f748.aa

MKFIIINLLSTIKIITFTVIVCLTILSIFQPIYILKENEISITRLGKIQRTENLAGLKYKIPLENVQIFPKIIL
 RWDGEPEQRIPTGGEEKQLIWIDTTARWKIADINKFYTTIKTMSRAYVRIDAAIEPAVRGVIAKYPLLEIIIRSSNDP
 IQLRLSNGILTPQETKINGIYKIKGKRKIIKEIIRIANNNTKDIGIEIVDVLIRKVTYDPSLIESVNNRMISERQQ
 IAEEQRSIGLAEKTEILGSIEKEKLKILSEAKATAAKIKAEGDREAAKIYSNAYGKNIIFYKFWQALESYKAVLKD
 KRKIFSTDMDFFQYLHKRN

TABLE 1. Nucleotide and Amino Acid Sequences

t748.aa

IFQPIVILKENEISTTRLGKIQRTENLPLKYKIPLIENVOIFPKIILRWGEPQRIPGEEKQLIWIDTTARW
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 IIEKEIIRIANMNTKDIGIEIVDVLIRK/T/DPSSLIESVNNRMISERQQIAEEQRSIGLAEKTEILGSIEKEKLKI
 LSEAKATAAKIKPREGDREAAKIVSNAYGPNIEFYKFWQALESYKAVLKDKRKIFSTDMDFFQYLHKRN

f748.nt

ATGAAATTATATAATTAATCTTTATTATCTACTATAAAGATTATAACCTTACAGTAATAGTTGCTTGACTATTT
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 AACTGAAATTAGCTGGACTTAAATATTAATACCATTAATTGAAAATGTGAAATATTCCAAAATCATTCTT
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 GATGGAATTGCAGACATAATAAATTTCACACAACAATAAAACAATGAGTAGAGCTTACGTTAGAATTGATGC
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t748.nt

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f764.aa

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 LIKAKDVEKDFQYEFNIYKQ

f764.aa

EKQFNZAIIFSDATEYFFEIQTPFIKNEILFINDKNLEIIKDKLKT
 KILLTHKSNEILNNEILKEKIFYLSKIKFSLKKSIDFLLNEKSIDLQKTLFRDKSLNNEDLEYLEKKGKEKNV
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 LIKAKDVEKDFQYEFNIYKQ

f764.nt

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 TTGAATAACAAATTAAATTGCAATAATTTCAGATGCAACTGAATTTTTTGAAATTCAAACAACTCCATT
 CATAAAACGAAACTATCTATTATAATGACAAATTAGAAATTATAAAAGACAAGCTAAAACAACAAAAAA

TABLE 1. Nucleotide and Amino Acid Sequences

ATACTATTAACCTATAAATCAAATAATGAAATTCTAAATAACGAAATTCTAAAAGAGAAAATTCTTATCTATCAA
 AAATAAAATTCTCTAAAAAAATCTATTGACTTCTGCTTAACGAAAATCAATAGATTGCAAAAACATTACT
 ATTTAGAGACAAATCTCTAAATAACGAAGACCTTGAATCTGGAAAAAAAAGGCAAAGAAAAAAATGTCAATATT
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 CTTTAAGAGATAATAATTATTATTTAAAAAGATACTAAATTGCGCTTTCTAAAAATATAAAATTGTATTAAAT
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t764.nt

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 GCAATACAAGAAAAGACTTAAATTATTAAGCTAAAATATAATCACCCCTAAAGAGCCTGATTTGATAAAAAT
 AGCAAAAGATGTTGAAAAGATTTCATATGAATTACATTATAACATAAACAATAA

f770.aa

MINFSKSFFYPLPIGKIFVLSGDMGSGKTSFLKGLALNLGISYFTSPTYNIVNVYDFINFKFYHIDLYRVSSLEEF
 ELVGGLEILMDLDSTIAIEWPQIALSIVPKDRLFSLTFKIVGSGRVVELNG

t770.aa

KTSFLKGLALNLGISYFTSPTYNIVNVYDFINFKFYHIDLYRVSSLEEFELVGGLEILMDLDSTIAIEWPQIALS
 IVPKDRLFSLTFKIVGSGRVVELNG

f770.nt

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 TTAA

t770.nt

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 TGTTGGGGATTGGAAATACCTTATGGATCTTGAACATTGCTATTGAATGCCACAAATTGCTTGTAGCATT
 GTTCCAAAGATAGATTATTTCTTAACTTTAAAATAGTAGGTTCAGGCAGGGTTGTAGAACTTAATGGTTAA

f790.aa

MNTKATTPLLLFLIQSLAFSSEIFEFKYIKGSKFRLEGTDNQKIYFNGHYNSSNTNIQISSEIKDIKENFASIK
 AFFRILKRENINEPYLLNEEFEEIFSVNKQGEYTIGANQKRPSVRGIPRFPKTPIKINEKWSYLAEEYIEASKIDK
 SIKDFVVKFNVNYEYKGKEEHNKGHYHILSNYESQNVKNISFYQKVDQKIYFDNEIGNTYKYSDFK
 NQHFKMIGNSLGRIVSIELPNDNLIEVENYIREKKIKAIeveKNNKGINLSFDIEFYPNSFQILQKEYKKIDLI
 AKLLEKFKKNNILIEGHTEQFGLEEEMHELSEKRAAIGNYLIKMKVKDKDQILFKGWGSQPKYPKSSPLKAKNR
 RVEITILNN

t790.aa

TABLE 1. Nucleotide and Amino Acid Sequences

SEIFEFKYIKGSKFRLEGTDNQKIQYFNGHYNSSNTNIQISSEIKDIKENFASIKAFFRILKRENINEPYLLNEEF
 EEIFSVNKQGEYTIGANQKRPNSVRGIPRFPKTPKINEKWSYLAEEYIEASKIDKSICKDFVVKFNVNVEYKGKEEH
 NGKHYHIILSNYESQYNVKNISFYQKVDQKIQYFDNEIGNTYKYSKDYIFEINQNNNQHFKMIGNSLGRIVSIELPN
 DNLIETEVENYIREKKIKAIeveKNNKGINLSFDIEFYPNSFQILQKEYKKIDLIAKLLEFKKNNILIEGHTEQF
 GLEEEEMHELSERAKRARAIGNYLIKMKVKDKDQILFKWGSGQPKYPKSSPLKAKNRREITILNN

f790.nt

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 TAATTCAAGCTCTAATACCAATATTCAAATTTCAAGTGAATAAAAGACATAAAAGAAAATTGCAAGCATTAAA
 GCTTTTTAGAATCTAAAAAGAGAAAATTAAATGAACCTTACCTATTAAATGAAGAGTTGAAGAAATCTTC
 GCGTAAATAAGCAAGGAGAATATACAATAGGAGCAAATCAAAAAGACCTCTGTTAGAGGTATTCAAGATTCCC
 AAAAACACCAATCAAATAATGAAAATGGTCATATCTTGAGAATATAGAACGCTAAAAATAGACAAA
 AGTATAAAAGATTCGTTGAAATTAAATGTTAATCTGAAATATAAGGCAAAGAGAGCACAATGGCAAGCATT
 ACCACATAATTCTTCGAATTATGAATCACAATACAATGTAAAAAACATCTTTCTATCAAAAAGTAGACCAAA
 AATTATTTGATAATGAAATTGCAATACATATAACAGCGATAAATATATATTGAAATAAATCAGAACAC
 ACCAACATTAAATGATTGAAACTCTCTGGCAGAATAGTTCAATTGAGCTTCCAAATGATAATCTTATTG
 AACTGAGGTTGAAAATTACATCCGAGAAAAAAATAAAAGCTATTGAAGTTGAAAAAAACAATAAAGGTATTAA
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 GCTAAACTCTGAAAATTAAAAAAATAACATAATAGAAGGACATACTGAGCAATTGGATTGAAAG
 AGATGCACGAGCTATCTGAAAAAGAGCTCGTCAATTGAAATTATTAATAAAAGTAAAGAAAAGACAAAGA
 CCAAATACTATTTAAAGGATGGGATCTCAAAAACAAAATATCCTAAGTCCTCCCCATTAAAGGCTAAAATAGG
 CGAGTAGAAATTACAATATTAATAACTAA

t790.nt

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 GAAGAAATCTCAGCGTAAATAAGCAAGGAGAATACAATAGGAGCAAATCAAAAAGACCTCTGTTAGAGGT
 TTCCAAGATTCCCAAAACACCAATCAAATAATGAAAATGGTCATATCTTGAGAAGATATAGAACGCT
 AAAATAGACAAAAGTATAAAAGATTCGTTGAAAATTAAATGTTAATCTGAAATATAAGGCAAAGAACGAC
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 GGATTGGAAGAAGAGATGCACGAGCTATCTGAAAAAGAGCTCGTCAATTGAAATTATTAATAAAAGTAAAG
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 GGCTAAAATAGCGAGTAGAAATTACAATATTAATAACTAA

f792.aa

MKIFIYWWVIVFFSVFKFSIYSLTDEEFFKKYSLFFVHKGFLSKNVNGKITKVQVNGINSRWVYPFYKLVPSRIT
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 KNLNRLIPQIYLGAGYYDIISAEFSKEETNNLYFSSGVYGDIFLISQKSGFIKKISFPFKQIIRILDLSKKNVE
 KILVRTYDNHFYSYINGQWVFIGKLSLQDQDFFEKSQRMQLAKNKGSIYLTAJLRLNKKAVDERFKFIKDSGMNAV
 VIDFKDDNGNLTYSSKLSLPNKLRSVKNFIDVPIYILKAKELGIYVIARCVFKDSKLYYDNFKHALWNKKTNP
 WAHLIKKVDSSGLVKYVQVEHWVDIFSPATWEYNISIAKEIQSGFVDEIQFDYIRFPSDGPVSLAISRMNKYEMQP
 VDALESFLIMAREQLYVPISVDIYGYNGWPTNSIGNQNIISMLSDYVDVISPMFYPSPHYTDDFLPSNFYYTKRAYRI
 YKEGSDRALAFSLDGVVIRPYVQAFLLGKERLVDDEIYLEYLKFQLGIKESFGSGFSLWNASNVYYMIK GSLKEY
 LDSF

TABLE 1. Nucleotide and Amino Acid Sequences

t792.aa

IYSLTDEEFFKKYSLFFVHKGFLSKNVNGKITKVQVNGINSRWVYPFYKLVPSRITSIYEDVYSSSFLTTSNNLY
 VSYDYSKNFRKLVGIDKFNSGAYITSSAFSQGDYKRIAIGTAIHINGIYLSVNGAISFKNLNRIPQIYLGAGYYDII
 SAIEFSKEETNNLYFSSGVYGDIFLISQKSGFIKKISFPFKKIIIRILDSSKNVEKILVRTYDNHFYSYINGQWV
 FIGKLSLQDQDFEKSQRMQLAKNKGSIYLTAYTLRNKAVDERFKFIKDSGMNAVVIDFKDDNGNLTYSSKLSLP
 NKLRSVKNFIDVPTYILKKAKELGIYVIARCVVFKDSKLYYDNFKHALWNKTKNPWAHLIKKVDSSGLVKYVQVE
 HWVDIFSPATWEYNISIAKEIQSGFVDEIQFDYIRFPSDGPVSLAISRMNKYEMQPVDalesFLIMAREQLYVPIS
 VDIYGYNGWFPTNSIGQNISMISLSDYDVISPMFYPSPHYTDDFLPSNFYYTKRAYRIYKEGSDRALAFSLDGVVIRP
 YVQAFLLGKERLVDDEIYLEYLKFLKGKESFGSGFSLWNASNVYYMIKGSLSKEYLDSF

f792.nt

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 AACCAAAGTTCAAGTCATGGATAAATTCTAGGTGGTTACCCCTTTATAAGCTTGTCTAGTCGAATTACT
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 AAAAATTTAAATCGTTGATTCCGCAGATTATTAGGTGCAGGATATTACGATATTAGTGCCTATTGAATTTT
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 TATAAAGAGGGGAGTGTGATAGAGCACTTGTCTTTCTTAGATGGGTTATTAGCCTTATGTTCAAGCTTTT
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 TTAGATTCTTTTA

t792.nt

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 AATAAGTTGAGATCTGTTAAAACCTTATTGATGTTCTTATATTCTTAAAGCAAAAGAGCTTGGAAATTATG
 TTATTGCTAGATGTTGTTATTAAAGATTCAAATTGATTATTATGATAATTAAACACGCCCTTGGAAATAA
 AAAACCAATAAACCTTGGCTCATTTGATTAAGATTCTAGTGGCTTGTGAAATATGTACAAGTAGAG

TABLE 1. Nucleotide and Amino Acid Sequences

CATTGGGTAGATATTTTCTCCTGCTACTGGGAATATAATATTCTATCGAAAAGAAATTCAATCTTGGAG
 TTGACGAGATACAATTGATTATATTAGATTTCATCAGATGGGCCTGTCCTGCAATCTCAAGAATGAATAA
 GTATGAGATGCAACCCGTTGATGCACTTGAATCTTTGATTATGCCAGAGAACAGCTTATGTTCTATTCT
 GTTGTATTTATGGTACAATGGCTGGTTCTACTAATAGTATTGGCAAGAGAACAGCTTATGTTCTATTCT
 TTGACGTCATATCTCTATGTTATCCTCGCATTATACTGATGATTTTGCCAAGCAATTTCACACAAA
 AAGAGCTTATAGGATTATAAAGAGGGAGTGATAGAGCACTGCTTTCTTAGATGGGTTGTTATTAGGCCT
 TATGTTCAAGCTTTTATTAGAAAAGAGATTGGTGGATGACGAGATTATGGAGTATTAAAGTTTCAGC
 TTAAAGGAATTAAAGAGTCATTGGTAGTGGCTTAGCCTTGAATGCATCTAATGTTATTATAATGATTAAGG
 TAGTTAAAAGAATATTAGATTCTTTAA

f797.aa

MSIKKFILTLIILSLAKNSFSENEINIFENENYIVKENIKTEIKKLQSFLASVDVAISQPYIELADLNGEPIKE
 LEGISYSFINVFSKIGSSAIISFDLSNEASKKYKIIKLEFLSPDKGNFINQLSSLTSGKQQSKKELAKDAYSFGL
 RTESLSKTIAEYYKDNNWYYILAAITVENNINKETEKYEIRINPKIYNDFQKKLRLHFKSNQIKKFPPIIE

t797.aa

KNSFSENEINIFENENYIVKENIKTEIKKLQSFLASVDVAISQPYIELADLNGEPIKELEGISYSFINVFSKIG
 SSAIISFDLSNEASKKYKIIKLEFLSPDKGNFINQLSSLTSGKQQSKKELAKDAYSFGLRTESLSKTIAEYYKD
 NWYYILAAITVENNINKETEKYEIRINPKIYNDFQKKLRLHFKSNQIKKFPPIIE

f797.nt

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 CAAACGAAGCTTCCAAGAAATACAAAATCATAAAATTAGAATTTTAAGTCCAGATAAAGGCAATTATTAACCA
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t797.nt

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 AACTGGTATTATATTAGCAGAATAACAGTAGAAAATAATAAAAGAAACTGAAAATACGAAATTAGAA
 TTAACCCAAAATATAATGATTTCAAAATTGAGATTACATTAAAAGCAACCAAATAAAATTTC
 AATACCCATTATAGAATAA

f799.aa

MKKHIIIGIIFVAILLFFKILLIPRIQNHENNKNNIKMIIISYKQDKNRLSLKINIKTKKTTNLGAKLDIYLDASKL
 IESNLLYISSKNFTTYANIIYQNESLLSIIILKSNGNNNVFSKRIKPRGKI

t799.aa

HENNKNNIKMIISYKQDKNRLSLKINIKTKKTTNLGAKLDIYLDASKLIESNLLYISSKNFTTYANIIYQNESLLS
 IIILKSNGNNNVFSKRIKPRGKI

TABLE 1. Nucleotide and Amino Acid Sequences

f799.nt

ATGAAAAAAACATATCAATTATTGGATAATCTTTGTGCAATTCTTTATTTTAAATTTATTAAATTCCAGAA
 TCTAAAATCACGAAAATAATAAATAAATCAAAATGATAATAAGCTACAAAGCAAGACAAAAACAGATTATCGCT
 AAAGATAAACATAAAACCAAAACTACCAACCTCGGAAAGCCAAACTAGATATTATCTAGACAGTAAATT
 ATTGAAAGCAATTGCTTATATAAGCAGCAAAACTTACAACATATGCTAATATAATCTATCAAATGAAAGTT
 TATTAAGTATAATATTAGAGTAATGCCAATATAATGTCTTTATAGTAAAAGAATAAAACCTAGAGGTTAAAT
 ATGA

t799.nt

CACGAAAATAATAAAATAATATCAAAATGATAATAAGCTACAAAGCAAGACAAAAACAGATTATCGCTAAAGATAAA
 ACATAAAAACAAAAAAACTACCAACCTCGGAAAGCCAAACTAGATATTATCTAGACAGTAAATTAAATTGAAAG
 CAATTGCTTATATAAGCAGCAAAACTTTACCATATGCTAATATAATCTATCAAATGAAAGTTATTAAGT
 ATAATATTAAAGAGTAAAGCAATAATAATGCTTTATAGTAAAAGAATAAAACCTAGAGGTTAAATATGA

f800.aa

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 DENGNIAIISIYSEGYIIYSYNEFSPLYKIVVNKNLLKTIDNQKKYNISIDKV
 FFEVNKKTLYVKTTYENIGDENINDLGIKIKDQYIYKMSLKKNKELEVINKIALPKNLLDDKQESFINIIKIQK
 DKIIIASTNMKNLSNNLWKLD SKGSIKEQIALIEPPNLMFLSESLSKDGI LSILYGGKTGVSVYWWNLNALLKL

t800.aa

KTNLNELGEEQFKIPFGTLPGAIMPLNNKFTNSKFDIKTYNGLVYIAEIKTNKLMIFNSYKLIQTYQNGIFKTNPD
 LKIKKIDFEGIQAIYPLKDFIIADKLNNKSKFNQKENIAYFMRILILNKNSSVEILQEGLNMPFPQIYDVNV
 DENGNIAIISIYSEGYIIYSYNEFSPLYKIVVNKNLLKTIDNQKKYNISIDKV
 FFEVNKKTLYVKTTYENIGDENINDLGIKIKDQYIYKMSLKKNKELEVINKIALPKNLLDDKQESFINIIKIQDKIIIASTNMKNLSNNLWKLD
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f800.nt

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 TTCAACTCATCGGAAAACTAACATACAAACATTCATAATGGAAATTAAACAAACCCGATTAAAAATAAAA
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 AAAATCAAATTCACCAAAAGAGAATTTCCTACTTCATGAGAATACTAATACAAACAAACTCATCTGTA
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 TTAAGAACAAATAGCTTAAATTGAGCCTCCAAATTAAATGTTCTCTGAGAGTTATCTAAAGATGAAACT
 TAGTATACTTATGGCGGAACACTGGTGTACTGTTACTGGTCGAATTAAATGCATTATTAAAATTATAA

t800.nt

AAAAACTTAAACGAATTAGGAGAAGAACATTAAACATTGGAAACACTTCCTGGTGCATAATGCCTCTGA
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 TAAATTAAATGATTTCACACTCATACGGAAACATACAAACATATCAAATGAAATTAAACAAACCCGAT
 TTAAAAATAAAAGATTTGAGGAACTCAAGCAATACCCACTAAAGATTATTGTCAGACA

TABLE 1. Nucleotide and Amino Acid Sequences

AACTAAATAATAAAAAATCAAACCAAAAAAGAGAATATTGCCTACTTCATGAGAATACTAATACTAAACAA
 AAACTCATCTGTAGAAATTGGGTCAAGAAGGTTAACCGGAATGCCATTCCACAAATTATGATGTTAATGTT
 GATGAAAATGCCAACATTGCAATAATATCAATATAAGCGAAGGATATATAATATATTCTTACAATAAGAATT
 CCCCGCTTATAAAATTACGTCAACAAAAACCTGTTAAAACAATAGACAATCAAAGAAAAATAACACATT
 AATAGATAAGGTTTTTGAGTCACACAAAAACTCTTATGTTAAACTACTATGAAACATTGGTGAC
 AATGAAAATATAACGATCTTGAATTAAAAGATCAATATATCTATAAAATGAGTTGAAAAAAACAAAG
 AATTAGAAGTGATAAATAAAATTGCTCTTCTAAACCTACTAGATGATAACAAAGAAAGCTTATAACATT
 AAAAATACAAAAAGACAAAATAATAGCATCTACTAATATGAAAATTATCTAACAATTAAATATGAAATTAGAC
 AGCAAGGGCTCAATTAAAGAACAAATAGCTTAATTGAGCCTCAAATTAAATGTTCTCTGAGAGTTATCTA
 AAGATGGAATACTTAGTATACTTATGGCGGAAACTGGTGTAGTGTACTGGTGAATTAAATGCATTATT
 AAAATTATAA

f810.aa

MYKLFLLFFIIFMFLSCDEKKSSKNLKSVKIGYVNWGGETAATNVLKVVFEKMGYNAEISVTTSIMYQYLASGKID
 GTVSSWVPTADKFYYEKLKTFVDLGANEGTIQGFVVPSSYVPISSISELKKGDKFKNMIGIDAGAGTQIVTEQ
 ALNYYGLSKEYELVPSSESVMLASLDSSIKRNEWILVPLWKPHAFSRYDIKFLDDPDLMGGIESVHTLVRGLE
 NDDFDAYYVFDHFYWSDDLILPLMDKNDKEPGKEYRAVEFVEKNKEIVKTWVPEKYKTLFD

t810.aa

CDEKKSSKNLKSVKIGYVNWGGETAATNVLKVVFEKMGYNAEISVTTSIMYQYLASGKIDGTVSSWVPTADKFYY
 EKLKTFVDLGANEGTIQGFVVPSSYVPISSISELKKGDKFKNMIGIDAGAGTQIVTEQALNYYGLSKEYELV
 PSSESVMLASLDSSIKRNEWILVPLWKPHAFSRYDIKFLDDPDLMGGIESVHTLVRGLENDDFDAYYVFDHFY
 WSDDLILPLMDKNDKEPGKEYRAVEFVEKNKEIVKTWVPEKYKTLFD

f810.nt

ATGTATAAATTATTTTATTATTTTATGTTTGTCTGTGATGAAAAAAAGAGTTCAAAGAATTAA
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 GTGCAAATTATGAAGGAACCATCAAGGTTTGTGGTGCAGCTATGTTCCAATTCCAGCATTAGTGAGCTTAA
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 GCGCTTAATTATTATGGATTAAAGTAAAGAGTATGAGCTAGTCCCTCAAGTGAGAGTGTATGTTGCAAGTTAG
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 GTGGGTTCCAGAAAATATAAGACCTTATTGATTAA

t810.nt

TGTGATGAAAAAAAGAGTTCAAAGAATTAAATCGTAAATTGGATATGTGAATTGGGGTGGAGAACCGCAG
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 AGCGATGATTAAATTGCCCCTTAATGGATAAAAATGATAAAGAGCAGGCAAAGAACACCGCAATGCGGTGAAT
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f814.aa

TABLE 1. Nucleotide and Amino Acid Sequences

MLVKRIVGKPIITMLILFSLLLMIISLYTFSRLKVDLLPGIDIPQISIHTVYPGASPREVEESRVLESGLSSVKNL
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 RYADEIIKPGLERLDGVAIVTVNGGSKKRVLIEVSQRLESYGLSLSRISIIIASQNLELSAGNILENNLEYLVEV
 SGKFKSIEEIGNVVIAYKIPDISSGINSPIEIKLKDIANIKTDFFEDLSEYVEYNGLPSISLSVQKRSDSNSIAVS
 NVVMNEIEKLKLSMPKDMKLEIASDSTDFIKASISTVVNSAYFGAMLAIFVIFFFLRSFRATIIIGISIPIAIVLT
 FCLMYFVNISLNIMSLAGLALGIGMVVDCSIVVIDNIYKYRQKGAKLISSSILGAQEMMLPITSSTFTSICVFGPF
 LIFKSELGVYGDFFKDFFTFTIVISLGVSLLVAIFLVPVLSSHYVGLYTSFQKNIKNAFIRKIDAFFASIYYFLEFL
 YINLLNIVLNHKLIFGLIVFFSFIGSLLLGLLDVTTFTRGKENSITINLNFPHKTNLEYAKFYSNRFLEIVKSEA
 KGKSIATLRADITFNVLFPLKEESRDNLTSQVDYDSIKYKIMNRIGNLYPEFNIEPSISGNALGGGDSIKIKI
 SANDFEYIKDYGKILVSMRKKEIPELVNPRLSISDFQLQIGVEIDRALVYNYGIDMNTILNELKANINGVVAQYV
 EKGLNYDIVLKLDLMDVKNLKDLEKIFITNSGVKIPFSSIATFEKTNKAESIYRENQALTIYLNAGISPPDNLTQ
 VTAKVVDIFINNKVPHKEGITLKVGEYNEFSNIMNQFKIIIMMAIIVVFGIMASQFESFLKPFIIIFTIPLTAIGV
 VLIHFLAGEKLSIFAAIGMLMLGVVVNTGIVLVDTGLLIKRGFGLREAIIESCRSRLRPILMSSLTSIIGLIPM
 AFSSSGSGNELLKPIAFTFIGGMTASTFLTFIPMLFEIFPTCFKFQI

t814.aa

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 PPIEIKLKDIANIKTDFFEDLSEYVEYNGLPSISLSVQKRSDSNSIAVSNVVMNEIEKLKLSMPKDMKLEIASDSTDF
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 SIVVIDNIYKYRQKGAKLISSSILGAQEMMLPITSSTFTSICVFGPFLIFKSELGVYGDFFKDFFTFTIVISLGVS
 LVAIFLVPVLSSHYVGLYTSFQKNIKNAFIRKIDAFFASIYYFLEFLYINLLNIVLNHKLIFGLIVFFSFIGSLL
 GLLLDVTTFTRGKENSITINLNFPHKTNLEYAKFYSNRFLEIVKSEAKGYKSIATLRADITFNVLFPLKEESRD
 NLTSQVDYDSIKYKIMNRIGNLYPEFNIEPSISGNALGGGDSIKIKISANDFEYIKDYGKILVSMRKKEIPELVNP
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 FSNIMNQFKIIIMMAIIVVFGIMASQFESFLKPFIIIFTIPLTAIGVVLIHFLAGEKLSIFAAIGMLMLGVVVNT
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 LFFIPMLFEIFPTCFKFQI

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 AGATATGCCATGAAATCATTAAACCTGGCCTGAAAGGCTTGATGGAGTTGCAATTGTTACTGTTAATGGTGGAA
 GTAAAAAGCGTGTAAATTGAAAGTTCTCAAACAGGCTGGAGTCTTATGGCTTCTTGTCAAGAATATCTC
 AATTATAGCATCCAAAATTGAACTTTCAGCTGGCAATATATTGAGAACAACTTGGAAATTGGTGAAGTT
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 TTGTTGACTGTTCAATTGTTGAAATTGAGAACAAATTGAAATTGAGCTTACCTTCTATTGTT
 TATTCTCGGAGCTCAGGAGATGTTGCCTATTACATCTCACTTCTATTGTT
 CTTATTCTCAAATCAGAACATTGGGTATATTGAGATTGTTCAAGACTTACATTACGATTGTT
 GTGTTCTCTTAAATGTTGCAATTGGTCTGTTATTCAAGCCACTATGTCGGTT
 AAAGAATATTAAGAATGCTTTATTAGGAAATCGATGCCTTTGCTAGTATT
 TTTAGAGTTTTG

TABLE 1. Nucleotide and Amino Acid Sequences

TATATCAATTATTAAATATAGTTAAATCACAAATTGATTGGGTTGATTGTTTTAGTTTATTGGCA
 GCTTGCTTTAGGATTATTGTTAGATGTGACAACCTTTACTAGAGGAAAGAGAACTCAATTACTATTAATTAAA
 TTTCCCCACAAAACAAATTGGAATATGCAAATTTATTCTAATAGATTAGAAATTGAAAAGTGAGGCT
 AAAGGATATAAAAGTATTATTGCTACTTGCCTGCTGATAGAATAACTTCACGTATTGTTCTCTCAAAGAAG
 AATCAAGAGATAATTAAACCCAAAGCGTAGATTACGATTCTATTAAATATAAAATTATGAATCGTATTGTAATCT
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 TTGTAATTCAGGCTTAGCATAAGTGAATTTCAGCTTAAAGGCTGAGATAGACAGAGCGCTAGTTATAA
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 GCATTTCAGCGGAAGTGGAAATGAACCTCTAAACCAATTGCAATTACTTTATTGGCGGAATGACAGCTAGCA
 CATTCTTACTTTGTTTATTCCATGCTTTGAAATTTCACATGTTCAAGTTCAAATCTAG

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 TTAATATTGAGCCTTCCATTAGTGGCAATGCTTGGTGGAGATTCTATTAAATTAAATT
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 ACATGAATACCATTAAATGAGTGAAGGCCAATTAAATGGTGTGTTGCTGGGAATATG
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 ATTCACTGAGTTAAATTCTTCTTCTAATAGCCACCTTGAAAAAACCAATAAGCGGAATCTATT
 GAGAAAATCAAGCTTAAACCATTTATCTTAATGCGGCTATTCTCCAGATGATAATT
 TAACCCAAGTAACCGCAAAAGTTGAGATTTTATTAAATAAGGTGCCCTAAAGGCAACTCTTAAGGTT
 GAAGGAGAATATAATGAA

TABLE 1. Nucleotide and Amino Acid Sequences

TTTCAAATATCATGAATCAGTTAAAATACTATTGATGGCTATTATTGTTGTTGGTATTATGGCTTCTC
 AATTGAACTTTTAAACCTTATTATTACAACTTAAACGGCAATAGGGGTTGTGCTTATACA
 TTTCTGCAGGAGAAAAGCTTCTATTGCTGCAATTGGTATGCTTATGCTTGTGGTGTGGTAAATACA
 GGAATTGTTCTGTAGACTATACTGGTTATTGATCAAGAGGGATTGGCTAAGAGAAAGCAATTATTGAATCTT
 GTCGTTCAAGGCTTAGGCCATTAAATGCTTCTTGACCTCAATAATAGGGTTATTCCAATGGCATTCTAG
 CGGAAGTGGAAATGAACCTCTAAAACCAATTGCATTACTTTATTGGCGGAATGACAGCTAGCACATTCTTACT
 TTGTTTTATTCCATGTTGAAATTTCACATGTTCAAGTTCAAATCTAG

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MLKNHSKLIQLKVMMIYLKKMGNDMTKFNYRIEIVSNLSLELDVFECIEKIEQELGESIYYSKIGNVYKGKK
 GEKHGNVWPEENFILIIYTSNQSIVERLKDIVDDLNRSYPTEGINLFVLRNS

t818.aa

KKMGNMDMTKFNYRIEIVSNLSLELDVFECIEKIEQELGESIYYSKIGNVYKGKKGEKHGNVWPEENFILIIYT
 SNQSIVERLKDIVDDLNRSYPTEGINLFVLRNS

f818.nt

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 GGAGAAAAGCATGGAATGGCCTTGGCCTGAAGAAAATTATTGATTATTACCTCCAATCAGTCTATTG
 TTGAGCGATTAAGGATAATTGTTGATGATTGAAATCGTTCTTACCCCTACAGAAGGGATTAAATCTTTGTTGAG
 AAATTCTAA

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AAGAAGATGGGAATGATGACTAAATTATAATTAGGATTGAAATAGTTCTAACTTATCTTAGAGCTTG
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 TCCAATCAGTCTATTGTTGAGCGATTAAGGATAATTGTTGATGATTGAAATCGTTCTTACCCCTACAGAAGGGATTAA
 ATCTTTTGTGAGAAATTCTAA

f820.aa

MLNNTYRIKTILTIFLAIITLLTIKYFTLMAFNNSPDNTISLKSNDIAKRGTIYDRNGKPIAFSSKSYSIGTNPQK
 IENIVSTSETLGAILQINSRILKEKLSNSNKGFLYIKRKIKREESDLIKRQAEGRLSNITLYPDYTRIYPRNTTS
 NITGFVGTDNLGLEGIEFSLNSILGKDCKTQQFLNEEPETNNIHLTIDMDIQQGVSKIAKKYFKENNPESLITLVM
 NSQNGEILSMVQFPQYDANFYSKYPEEIRKNLSSSLTYEPEGSINKIFTVAIILESGKLNLEEKFLDNGIYQKQFPS
 GEKITIKTLNPPYKHIDSTEILLYSSNVGIAYITEKVSNEYFYKLLDFGFGEKVGVPFPGETKGLLNHYSKWSGR
 SKATIGFGQEIGVSAVQILQAAISLNNGIMLKPRIKKISNDKGENIKEFDKEEIRKVISKNSAQKVLKMMREVV
 NKGGIPNLKIKNLDisAKSGTSQAI DRKTGKYSEEDYTSSILAIYPTEQPKIYIYVYRPPKIIYGTriaAPMAK
 EIIIEFIEHQNTIAYKKIKMPSKIKIPKAETNYKNKTYLPNFINLSKREAIIDLKYYKNTMKIKINGDGFVYKQSI
 SPNTKLEDITELELYLK

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FNNSPDNTISLKSNDIAKRGTIYDRNGKPIAFSSKSYSIGTNPQKIEIVSTSETLGAILQINSRILKEKLSNSNKG
 FLYIKRKIKREESDLIKRQAEGRLSNITLYPDYTRIYPRNTTSNITGFVGTDNLGLEGIEFSLNSILGKDCKTQ
 QFLNEEPETNNIHLTIDMDIQQGVSKIAKKYFKENNPESLITLVMNSQNGEILSMVQFPQYDANFYSKYPEEIRKN
 LSSSLTYEPEGSINKIFTVAIILESGKLNLEEKFLDNGIYQKQFPSGEKITIKTLNPPYKHIDSTEILLYSSNVGIA
 YITEKVSNEYFYKLLDFGFGEKVGVPFPGETKGLLNHYSKWSGRSKATIGFGQEIGVSAVQILQAAISLNNGIM
 LKPRIKKISNDKGENIKEFDKEEIRKVISKNSAQKVLKMMREVVNKGIPNLKIKNLDisAKSGTSQAI DRKTGK

TABLE 1. Nucleotide and Amino Acid Sequences

YSEEDYTSSILAIYPTEQPKYIIYIVYRYPKIIYGYTRIAAPMAKEIIEFIEHQQNTIAYKKIKMPSKIKIPKAET
NYKNKTYLPNFINLSKREAIIDLKYYKNTKIKINGDGFVYKQSISPNTKLEDITELYLK

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ATTCACACTAATGGCCTTCATAACAGCCAGACAACACAATATCTTAAAGTCAAATGATATTGCCAAAAGAGG
ACAATTTATGATAGAAATGGCAAACCAATAGCATTCTCTCAAATCCTACTCAATTGGTACAAATCTCAAAAA
ATAGAAAATATTGTAAGCACATCTGAAACTCTTGGTCAATACTTCAAATTAAATTCAAGAATTAAAGGAAAAGC
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AGCTGAAGGCAGGCTTCAACATCACTTATATCCTGATTACACAAGAATTATCCCTTCAGGAATACCAACAGC
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GCAATAATGGAATAATGCTAAAACCTAGAATAATAAAAATAAGCAACGATAAGGAGAAAATATTAAAGAATT
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TCCCTAAAGCTGAAACTAATTACAAAACATACTTACCAAATTTCATCAACCTTCTAAAAGAGAAGCAAT
AGACATACTAAAATACTATAAAATACATGAAAATTAAATGGCAGTGGATTGTTACAAGCAAGTATA
TCCCCAATACAAATTAGAAGATATAACAGAGCTGAACTGTATTAAATAA

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CAAACATCACTTATATCCTGATTACACAAGAATTATCCCTTCAGGAATACCACAAGCAATATTACTGGTTTGT
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CTAAAACCTAGAATAATAAAAATAAGCAACGATAAGGAGAAAATATTAAAGAATTGATGAGAGAAGTTGTA
GAAAAGTAATATCCAAAATTCAAGCACAAAAGTTAAAATGAGGAGAAGTGTAAATAAAGGTGGAATTCC
AAATCTTAAATTAAATCTTGTACATTCTGCAAAAAGTGGACATCTCAAGCTATTGATAGAAAACGGAAA
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TATACAGATAACCAAAAAATAATACGGAACAAAGAATAGCAGCCCAATGGCAAAGAAATAATAGAATT
TGAGCACCAACAAAATAACATAGCATATAAAAATTAAATGCCATCAAATCAAGATCCCTAAAGCTGAAACT
AATTACAAAACACATACTTACCAAATTTCATCAACCTTCTAAAAGAGAAGCAATAGACATACTAAAATAC

TABLE 1. Nucleotide and Amino Acid Sequences

ATAAAAAATACTATGAAAATAAAATGGCGATGGATTGTTACAAGCAAAGTATATCCCCAATACAAAATT
AGAAGATATAACAGAGCTTGAACGTGATTAAAATAA

f831.aa

MAKNNLLVFFIAIIIFVFVSIIVVFYNSLGKDYVKSGGEIVENLEKDLNDYLKENDAKEREKIFLRIRELISKEKEI
SSYFISRFYLARAVYFQSQAQYDEAIKDLDIVIKAKGIESEIAFLNKAAYEKMLKEDALLVYEDLINSTSLDFL
KVRALLSKAILIEEKDKELAVKVYEEIVKF PYENNLYINMANNKILELKQN

t831.aa

YNSLGKDYVKSGGEIVENLEKDLNDYLKENDAKEREKIFLRIRELISKEKEISSYFISRFYLARAVYFQSQAQYDE
AIKDLDIVIKAKGIESEIAFLNKAAYEKMLKEDALLVYEDLINSTSLDFLKVRA
LLSKAILIEEKDKELAVKVYEEIVKF PYENNLYINMANNKILELKQN

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TTAA

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GAAGAGATTGTTAAGTTCCGTATGAAAATAATTATATAATGGCAAATAATAAAATTAGAACTTAAGC
AAAATTAA

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MKAIGNAILLNMPLISIGISIGVARMGQGTAAALGGGLIGYLTFNITENYFIEAFSGLVEAETMSSVGRINFFGVQT
LNTGIAGSLAVGLLVGYLHNKFYNMKLPKPFVFFSECHFVPIVIIILPFCVFLAIFFCCLIWSSFDDLIASLGLFVFR
FEYFGSFLYGFNRLPLGLHSILSFPFEFTSLGGVEIVNGDTVRGLKNIFYAQLLDPSLGKFSSGFAKISSGFY
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VTIGNSFSTGFLDFMFGLQGNSKTNWISVPLGAMFFALYYFTFSWLYRYFDFQIFVTDPPFFEGQEGKLESLG
IAHLLIQGLGGFDNITKLDVCSTRLHVDVNTTELVDNNLLKEAGVLKIGLVNGKVQLFYGSNVYYIKNAIDTYS
PKSLFEASVMAVDNVKKGFKTYIEMKEDKLEKQGKSGKTYKLSELEED

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KLPKPFVFFSECHFVPIVIIILPFCVFLAIFFCCLIWSSFDDLIASLGLFVFRFEYFGSFLYGFNRLPLGLHSIL
SFPFEFTSLGGVEIVNGDTVRGLKNIFYAQLLDPSLGKFSSGFAKISSGFYLSIMFGLPGAALGVYKGIVHEDKNK
VAALLFSGALTAFLTGITEPLEFLFIFTAPLLYFVHAAYSGFALLNFFNVTIGNSFSTGFLDFMFGLQGNSK
TNWISVPLGAMFFALYYFTFSWLYRYFDFQIFVTDPPFFEGQEGKLESLGIAHLLIQGLGGFDNITKLDVCSTRL

TABLE 1. Nucleotide and Amino Acid Sequences

HVDVVNTELVDNNLLKEAGVLKIGLVNGKVQLFYGSNVYYIKNAIDTYSPKSLFEASVMVAVDNVKKGFKYIEMK
EDKKLEKQGKSGKTYKLSELED

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TGAGGCTTTTCAGGGCTTGTGAAGCAGAGACAATGCTTCGTGCGTATAAATTGGTGTCAA
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AGCTACCCAAACCTTTGTGTTTTCAAGAGTGCCTATAGTAATAATTACCCCTTGTGTT
TTGGCTATATTTTGTTGATTTGGTCAAGTTGACGATTAATTGCATCTTAGGTTGTTGTTAGG
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TATTGCTCAGCTATTAGACCCATCACTGGTAAATTTCATCAGGCTTGCCAAAATTAGCAGTGGATT
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ATTGCGCATTTTAATTCAAGGCTTGGGATTGATAATTACAAAGCTGATGTTGTTCTACAAGATTG
CATGTAGATGTTGTTAACTGAGCTTGTGATAATTGCTTAAAGAGGCTGGAGTTCTAAAATAGGGCTG
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GAGTCTTTGAGCTAGTGTATGGTGCAGTTGATAATTGAAAGGTTAAAACCTATTGAAATGAAAG
GAAGACAAAAAAACTTGAAAAGCAAGGTAATCAGGAAAAACCTATAAGCTAGCGAATTAGAAGAAGATTAG

f850.aa

MRFKKIFLIFIISNLKVSYNYAIQYKNEGIDKYYFEILNDGFGFSLSDFFDDLRSGLIFTYVSKYNFIINLEA
HMLTYRGYKDSPKSLISRTDLIEIGFMYFPILLINGKNGEIDLIGVKNLLFGDWGGHLMQSIIHLILNQH
IPSIKSYDSNYRGFLSFALNYSYMNFLNLENYMDLSYFADYFIKNSIGITLKNENIGFDIKLYSQIQNQIKSLKT
YSKTQEAETGIGIINYQFYSKNFFITNNLNKNFSTKENFLSVGGFGIIITPEEYKKKISESNNEFNVISNNFYFGFD
IMIPLKIRNSLFYKINENINHYFSISTNYTNYNETNSFTNQLSSGIMYEFLPQKTFNPYLIISGLFFAYQNQNNKDI
KSISRPIRIKNILQVGIENELGFLFKMLKRNTEYIFKISKVNYIPIAYNLDEKKLEKHSINFNYLGIGIVVK

TABLE 1. Nucleotide and Amino Acid Sequences

t850.aa

YSNYAIQYKNEGIDKYYFEILNDGFGFSLSDFFDDLRSGLIFTYVSKYNFIINLEAHMLTYRGYKDSPKSLISR
 TDLIEIGFMYYPILLINGKNFGEIDLIGVKNLLFGDWGGHLMQSIIHLILNQHRIPIPSIKSYDSNYRGFLSF
 ALNYSYMFLNLENYMDLSYFADYFIKNSIGITLKENENIGFDIKLYSQIQNQIKSLKTYSKTQEATGIGINYQFY
 SKNFFITNNLNKNFSTKENFLSVGGFGIIITPEEYKKISESNEFNVISNNFYFGFDIMIPLKIRNSLFYKINEN
 INHYFSISTNYYTNYNETNSFTNQLSSGIMYEFLPQKTFNPYLIISGLFFAYQNQNNKDIKSISRPIRIKNILQVGIE
 NELGFLFKMLKRNTEYIFKIVSKVNYIPIAYNLDEKKLEKHSINFNYLGIGIVVK

f850.nt

ATGCGGTTAAAAAAATTTTAATAATATTATAATTCTAATTAAAAGTTTATTCTTATAATTATGCAATCC
 AATATAAAAATGAAGGTATTGACAATATTATTTGAAAATCTAAATGATGGATTGGATTTCTTATAAGCGATT
 TTTGATGACTTGAGAAGTGGTCTCTTATTTCACCTATGTTCAAAATACAATTTTATAATAAAATTAGAAGCA
 CACATGTTAACCTATAGGGTTATAAGACTCTCCGAAATCTTAAATTAGTAGAACAGACTTAATTGAAATAGGCT
 TCATGTAATTTCACCTATGCTAATTGAAAGTGGGAGGGCATTAAATGCAAAGCATAATTCACCTCATTTAAATCAACACCGTCCA
 ATTCCAAGTATAAAAGCTACGACAGCTACAATTAGAGGATTTAAGCTTTGCTCTAAATTACTCTACATGA
 ATTGTTAAATTAGAAAATTATGGACTTATCTTATTTCAGATTATTTATTAAAACAGTATTGAAATTAC
 CTTAAAAAATGAAAATTGGATTGATATAAAACTTATTCCAAATTCAAATCAAACGCTCAAACAA
 TATTCAAAAACACAAGAACAGGAATTGGAATAAATTCAATTACTCTAAATTGTTACATGTTAAATTTCATAACCA
 ATAATTAAACATTAAAAATTTCACCAAAAGAAAATTCTTAAGCTGGGATTTGGAATAATCATTACACC
 TGAAGAATACAAAAAAATCAGAATCAAATAATGAATTAAATGTTATAAGTAATAATTTCAGTTGGATTGAT
 ATTATGATCCCATTAAAATAAGAAATTCAATTGTTATAAAATAATGAAAACATCAACCATTACTTTCAATAT
 CAACAAATTACTAAATTATAATGAAACTAATAGCTTACAAATCAATTATCATCAGGCATCATGTATGAATT
 TTTACACAAAAACATTCAATTCTTACCTAATTCTGGATTATTTTGCCTATAATCAAACAAATAAGATATC
 AAAAGCATCTCAAGACCAATAAGAATAAAAACATTCTCAAGTTGAAATTGAAATTAGGATTTGTTCA
 AAATGCTAAAATACCGCAACACTGAGTATATTCTAAAATATTCAAAGTTAACTATATTCTATAGCTTAA
 CTTAGATGAAAAAAATTAGAAAACATTCTATTAAACTTTAATTAGGAATTGGAATTGCTTAAATAA

t850.nt

TATTCTTATAATTATGCAATCCAATATAAAATGAAGGTATTGACAATATTATTTGAAATACTAAATGATGGAT
 TCGGATTTCTTAAAGCATTGATGACTTGAGAAGTGGTCTCTTATTTCACCTATGTTCAAATACAA
 TTTTATAATAAAATTAGAACACATGTTAACCTATAGGGTTATAAGACTCTCCGAAATCTTAAATTAGTAGA
 ACAGACTTAAATTGAAATAGGCTCATGTAATTCTTAAATTGAAATGAAAACATCAACCATTACTTTCAATAT
 TAGACTTGGGAATTGGAGTTAAAACCTTATTGAGACTGGGAGGGCATTAAATGCAAAGCATAATTACCT
 CATTAAATCAACACCGTCCAATTCAAGTATAAAAGCTACGACAGCTACAATTATAGAGGATTTAAGCTTT
 GCTCTAAATTACTCTTACATGAATTCTTAAATTGAAATTATGGACTTATCTTATTTCAGATTATTTA
 TTAACAGTATTGAAATTACCTTAAATTGAAATTGATATAAAACTTATTCCAAATTCAA
 TCAAATCAAAGCCTAAAACATATTCAAACAAACACAAGAACAGGAATTGGAATAAATTCAATTTCAC
 TCTAAAATTCTTCAATTACCAATAATTAAACATTAAACATTCTTCAACCAAGAAAATTCTTAAAGCTGGGG
 GATTGGAATAATCATTACACCTGAAGAATACAAAAAAATCAGAATCAAATAATGAATTAAATGTTATAAGTAA
 TAATTCTTACTTGGATTGATATTGATCCCATTAAAATAAGAAATTCAATTCTTATAAAATAATGAAAC
 ATCAACCAATTACTTCAATTCAACAAATTACTAAATTGAAACTAATAGCTTACAAATCAATT
 CATCAGGCATCATGTAATTCTTACACCTGAAGAATACAAAAAAATCAGAATCAAATAATGAATTAAATGTTATAAGTAA
 TAATCAAACAAATAAGATATCAAAGCATCTCAAGACCAATAAGAATAAAAACATTCTCAAGTTGAAATTGAA
 AATGAATTAGGATTTGTCAAAATGCTAAAATACCGCAACACTGAGTATATTCTAAAATATTCAAAGTTA
 ACTATATTCTTACAGCTTAAACTTAGATGAAAAAAATTAGAAAACATTCTATTAACTTTAATTATTAGGAAT
 TCGGAATTGCTTAAATAA

f853.aa

MKSFLFWVILGTVGSSFAQNTVAIINLYKNEIITKTGFDKVDIFKKTQGRDLTDAEKKQVLQVLIAVDLFSQE
 ASKQGIKISDDEVMQTIIRTQFGLVNFDEQIKQMIEKQGNTWGELLSSMKRSLSQQQLVKQAOQPKFSEIKTPSEK
 EIVEYYEANKTKFVNPDISRVSHIFFSTDKKRSVDLDQAKNILSQIRSKITFEEAVRKYSNDESSAKNGDLGF

TABLE 1. Nucleotide and Amino Acid Sequences

LSRGDQNAQNLGADFVKEVFNFKGDISSPIASKEGFHIVKVTEKYAQRFLGLNDKVSPTADLIVKDAIRNNMIN
VQQQQIVVQVQQDMYGKLNKSANIQILDSSLK

t853.aa

QNTPVAIINLYKNEIITKTGFDSKVDIFKKTQGRDLTDAEKQVLQVLIAADVLFSQEASKQGKISDDEVMQTIRT
QFGLVNFTEQIKQMIKEQKGTNWGELLSSMKRSLSQKLVLKQAQPKFSEIKTPSEKEIVEYYEANKTKFVNPDIS
RVSHIFFSTKDKKRSVLDQAKNILSQIRSKKITFEEAVRKYSNDESSKAKNGDILGFLSRGDQNAQNLGADFVKE
VFNFNFKGDISSPIASKEGFHIVKVTEKYAQRFLGLNDKVSPTADLIVKDAIRNNMINVQQQIVVQVQQDMYGKLN
KSANIQILDSSLK

f853.nt

ATGAAGAGTTTTATTTGGTAATATTGGGAACTGTAGGGATTAGCTTTGCTAAAATACTCCTGTTGCTA
TTATTAAATTATATAAGAATGAAATTATTACTAAAACGGTTGATTCTAAGGTTGATATTTAAAGACCCA
AGGTAGAGACTTAACGTGAGAAAAGCAAGTTCTGCAAGTTAAATAGCAGATGTTAGTCAAGAG
GCTTCAAAGCAAGGAATTAAATCTCAGATGATGAGGTTATGCAAACAATTAGAACATTAGAACATTGGCTTGGAATT
TTACTGATGAACAAATCAAGCAAATGATAGAAAAACAGTACAATTGGGGCGAGCTTGTCTCAAGTAAAG
ATCTCTGTTCTCAAAGCTGTTAAAGCAAGCTCAGCCTAAGTTCTGAAATTAAACCTCTAGTGAAGAAA
GAAATTGTTGAGTATTATGAGGCTAATAAAACTAAGTTGAAATCCCAGATATTCAAGAGTTAGTCATATCTTT
TTTCTACTAAAGATAAAAAAGATCAGATGTTAGATCAAGCAAAATATTAAAGCCAATAAGATCAAAAAAA
AATTACTTTGAAGAAGCTGTAAGAAAATATTCAAATGACGAATCTCTAAGGCTAAAATGGTGTCTGGGTT
TTATCAAGAGGTGATCAAATGCTCAAATCTCTGGAGCCGATTTGTGAAAGAGGTTTAAATTAAAGG
GTGATATATCTCGCTATTGCTCAAAGGAAGGGTTCATATTGTTAAAGTTACAGAAAATATGCTCAGAGATT
TTAGGTTGAATGATAAAGTGTCTCCTACTGCAGATTGATGTCAGGAAATGCAATAAGAAATAACATGATTAAT
GTTCAACACAGCAAATTGTTCAAGTACAGCAAGATATGTATGGTAAGCTAACAGTCTGCAAATATACAAA
TCTGGATTCTAGTCTAAATAA

t853.nt

CAAAATACTCCTGTTGCTATTAAATTATATAAGAATGAAATTATTACTAAAACGGTTTGTCTAAGGTTG
ATATATTAAAGACCCAAGGTAGAGACTTAACGTGAGAAAAGCAAGTTCTGCAAGTTAAATAGCAGA
TGTTCTTTAGTCAGAGGCTCAAAGCAAGGAATTAAATCTCAGATGATGAGGTTATGCAAACAATTAGAACT
CAATTGGGCTTGTGAATTAACTGATGAACAAATCAAGCAAATGATAGAAAACAAGGTACAATTGGGGCGAGC
TTTGTCTCAATGAAAAGATCTCTGTTCTCAAAGCTGTTAAAGCAAGCTCAGCCTAAGTTCTGAAAT
TAAAACCTCTAGTGAGAAAGAAATTGTTGAGTATTATGAGGCTAATAAAACTAAGTTGTTAAATCCCAGATATTCA
AGAGTTAGTCATATCTTTTACTAAAGATAAAAAAGATCAGATGTTAGATCAAGCAAAATATTAA
GCCAAATAAGATCAAAAAAAATTACTTTGAAGAAGCTGTAAGAAAATATTCAAATGACGAATCTCTAGGCTAA
AAATGGTGTCTGGGTTTATCAAGAGGTGATCAAATGCTCAAATCTCTGGAGCCGATTTGTGAAAGAG
GTTTTAATTAAATAAGGGTGTATATCTCGCTATTGCTCAAAGGAAGGGTTCATATTGTTAAAGTTACAG
AAAAATATGCTCAGAGATTAGGTTGAATGATAAAGTGTCTCCTACTGCAGATTGATGTCAGGAAATGCAAT
AAGAAATAACATGATTAATGTTCAACACAGCAAATTGTTCAAGTACAGCAAGATATGTATGGTAAGCTAAC
AAGTCTGCAAATATACAAATCTGGATTCTAGTCTAAATAA

f859.aa

MKLPKLYKLILLFLFTTRLFSVKDEKSDNKLLELFNSVETKIKKNSKNYDSNSNSKIKKESILKRDINSEKNINSN
IYIQLSKKINYPNRNLGNNINQKTANDVNFTKTSYVVKVYPNYKDDNFQEIKNANKFPAKTEKTHMLIGPILKDNLG
IIKMLKTKGYTLIEYIEDNN

t859.aa

VKDEKSDNKLLELFNSVETKIKKNSKNYDSNSNSKIKKESILKRDINSEKNINSNIYIQLSKKINYPNPNLGNIN
QKTA

f859.nt

TABLE 1. Nucleotide and Amino Acid Sequences

ATGAAATTACCAAAACTTACAAATTAAATCACTACTCTTCTTTACAACAAGATTGTTCAAGTAAAAGATGAAA
 AATCAGACAATAATGGAATTATTTCAACGAGAAACAAAAATCAAAAAAATTCTAAAATTACGACTCAAA
 TTCAAAACAGCAAAAGATCAAAAAAGAATCAATTAAAAGAGATAACAAACAGCGAAAAAAATATAAATCCAAT
 ATATACATACAAAATCAAAAAAATTAAATTACCCCAACAGAAATTAGGCAATAATATCAATCAAAAATGCAA
 ATGATGTAATTTCACAAAATGTTAGTTATGCCAATCTAAAGACGATAACTTCAGAAATTAA
 AAATGCTAATAAATTCCAGCTAAAACCGAAAAACTCACATGCTAATCGGCCAATATTAAAGATAATCTAGGA
 ATAATAATTAAATGCTAAAACAAAGGGATAACTTAATAGAATACATAGAGGACAATAATTAA

t859.nt

GTAAAAGATGAAAATCAGACAATAATTGGAATTATTTCAACGAGAAACAAAAATCAAAAAAATTCTAAA
 ATTACGACTCAAATTCAACAGCAAAAGATCAAAAAAGAATCAATTAAAAGAGATAACAAACAGCGAAAAAA
 TATAAATTCAATATATAACATACAAAATCAAAAAAATTAAATTACCCCAACAGAAATTAGGCAATAATATCAAT
 CAAAAAACTGCAATGATGTAATTTCACAAAATGTTAGTTATGCCAATCTAAAGACGATAACT
 TTCAAGAAATTAAAATGCTAATAAATTCCAGCTAAAACCGAAAAACTCACATGCTAATCGGCCAATATTAA
 AGATAATCTAGGAAATAATTAAATGCTAAAACAAAGGGATAACTTAATAGAATACATAGAGGACAATAATTAA

f861.aa

MKNFKEVIIIFDSGIGGLSYFKYIKSRIGGCQYVYVADNKNFPYGEKSPEYLLEAVLFLIEKLKKIYNIGALVLAC
 NTISVSYNKLNFVFPVYTLPDVSSVSDLVLKRVLLIATNTTLESKFVQDVNIHNDLIVKAAGELVNFVEYGEN
 YKKYALRCLEALKFEVNTGREIVFLGCTHYLHLKVMIEDFLKIPVYENRELVVKNLIRSMNFSEHKGNYYKNDFD
 FVDEFYLTKNLTQNFCKKYNLRFKGMIV

t861.aa

RIGGCQYVYVADNKNFPYGEKSPEYLLEAVLFLIEKLKKIYNIGALVLACNTISVSYNKLNFVFPVYTLPDVSS
 VSDLVLKRVLLIATNTTLESKFVQDVNIHNDLIVKAAGELVNFVEYGENYKKYALRCLEALKFEVNTGREIVFL
 GCTHYLHLKVMIEDFLKIPVYENRELVVKNLIRSMNFSEHKGNYYKNDFDVDEFYLTKNLTQNFCKKYNLRFKGMIV

f861.nt

ATGAAAATTCACAAAGTAATAATTATTTGATTGAGGAATAGGAGGGCTTCTTATTAAATATATTAAA
 GTAGAATAGGGGATGCCAATATGTTATGTTGCCGATAATAAAATTCCCTTATGGAGAAAAAGTCCTGAATA
 TCTCTAGAACGAGTTGTTGATTGAGAGCTTAAAAAATCTATAATTGGTCATTAGTTGGCTTGT
 AATACAATTCTGTTAGTGTATACAATAAAATTAAATTGTTCCAGTAGTCATAACTTGCAGATGTAAGTT
 CAGTTTCAGATCTGTTAAAAAGAGTTCTTGTGATTGCAACAAATACTACTCTGAAAGCAAATTGTTAAGGA
 TCAAGTAAATATACATAATGATTGATTGAAAAGCTGCTGGAGAGCTGTTAATTGTTGAATATGGAGAGAAT
 TACAAAAAATATGCTCTAGATGTTAGAGCTTAAATTGAGTTGTAATAACTGGTAGAGAAATTGTTTTC
 TTGGATGCACGCATTATTGCACTTAAGGTAATGATAGAAGATTTTAAATTCTGTTATGAGAATCGTGA
 ATTAGTGGTAAAAAATCTTATTAGATCAATGAATTCTGAAACACAAAGTAATTATTATAAGAATGATTGAT
 TTTGTAGATGAGTTATTGACCGAAAATAAAATTGACTTTATCAAAATTGCAAAATATAATC
 TTCGCTTAAGGAATGATAGTTGA

t861.nt

AGAATAGGGGATGCCAATATGTTATGTTGCCGATAATAAAATTCCCTTATGGAGAAAAAGTCCTGAATATC
 TTCTAGAACGAGTTGTTGATTGAGAGCTTAAAAAATCTATAATTGGTCATTAGTTGGCTTGTAA
 TACAATTCTGTTAGTGTATACAATAAAATTAAATTGTTCCAGTAGTCATACTTGCAGATGTAAGTTCA
 GTTTCAGATCTGTTAAAAAGAGTTCTTGTGATTGCAACAAATACTACTCTGAAAGCAAATTGTTAAGGATC
 AAGTAAATATACATAATGATTGATTGAAAAGCTGCTGGAGAGCTGTTAATTGTTGAATATGGAGAGAATTA
 CAAAAAATATGCTCTAGATGTTAGAAGCTTAAATTGAGTTGTAATACTGGTAGAGAAATTGTTTCTT
 GGATGCACGCATTATTGCACTTAAGGTAATGATAGAAGATTGTTAAAAATTCTGTTATGAGAATCGTGAAT

TABLE 1. Nucleotide and Amino Acid Sequences

TAGGGTAAAAATCTTATTAGATCAATGAATTCTGAACACAAAGGTATTATAAGAATGATTTGATTT
 TGTAGATGATGAGTTTATTCGACGAAAATTTGACTTTATCAAAATTGCAAAAATATAATCTT
 CGCTTAAAGGAATGATAGTTGA

f363.aa

MIRLKVLILCLFGIFVLNGFADTNFEFNFGGGVAFPVSPFSSFYNEALEINAKLKQNLPSDLSPIEKEEIVQNFSD
 LANIAGIRYGTYAQFGAKFDDFVSIGFELLFNINLLKAIKRSRGTANENFSFIMAITPRFYTKLDFFVLA
 MGPKINIATSSADSVLAELGTMGWDIGARLSFSFILEGYVWNINKNPKFSDFKFGIGFEFGIV

t363.aa

DTNFEFNFGGGVAFPVSPFSSFYNEALEINAKLKQNLPSDLSPIEKEEIVQNFSDLANIAGIRYGTYAQFGAKF
 DDFVSIGFELLFNINLLKAIKRSRGTANENFSFIMAITPRFYTKLDFFVLA
 MGWDIGARLSFSFILEGYVWNINKNPKFSDFKFGIGFEFGIV

f363.nt

ATGATTAGGCTTAAAGTTAAATTGTGTTATTGGGATTTGTGTTAAATGGTTGCAGATACTAATTG
 AATTCAATTGGTGGTGGGTTGCTTTCTGTTAGTCCCTTTCAAGCTTTACAATGAGGCTTAGAGATTAA
 TGCAAAGCTTAAGCAAAATTGCCCTCAGATTATCCCCAATAGAAAAAGAAGAGATAGTCAA
 TTGCAATATTGCTAAAGCTGGAATAAGATATGGAACCTACGCTCAATTGGCGCTAAATTGATGATTGTT
 CTATTGGATTGAGCTTTGTTAACATTAATCTCTAAAGCAATAAACGCTTCGATGGAAC
 ACAGGTCTAACAGATCAATATAGCGACTTCTGCGGATTCTGTTAGCAGAACTGGAA
 ATGGTCTAGACTTCATTCTTTAATTCTGAAGGGTACTATGTTGGAATATTAAA
 TGATTCAAGTTGGAATAGGTTGAAATTG
 GAATTGTGTAG

t363.nt

GATACTAATTGAAATTCAATTGGTGGTGGGTTGCTTCTGTTAGTCCCTTTCAAGCTTTACAATGAGG
 CTTAGAGATTAATGCAAAGCTTAAGCAAAATTGCCCTCAGATTATCCCCAATAGAAAAAGAAGAGATAGTCCA
 AAATTGATTTCCGATTAGCAATATTGCTAAAGCTGGAATAAGATATGGAACCTACGCTCAATTGGCGCTAAATT
 GATGATTGTTCTATTGGATTGAGCTTTGTTAACATTAATCTCTAAAGCAATAAGCTTCGATGGA
 CTGCAAATGAAAATTCTCGTTATTATGCAATAACACCAAGATTATACAAAATTAGATTGTTAGC
 TTAGCGTTTCACAGGCTCTAACAGATCAATATAGCGACTTCTGCGGATTCTGTTAGCAGAACTGGAA
 ATGGGCTGGGATATTGGTGCTAGACTTCATTCTTTAATTCTGAAGGGTACTATGTTGGAATATTAAA
 ACCCTAAATTCTGATTCAAGTTGGAATAGGTTGAAATTGGAATTGTGTAG

f368.aa

MIDLQEKQEILIKNKFRAKVFGLMSIGLLISAVFAYATSENQTIKAIIFNSMSFMAMILIQFGLVYAIISGALNK
 ISSNTATALFLLYSALTGVTLSSIFMIYTQGSIVFTFGITAGTFLGMSVYGYTTTDLTKMGSYLI
 MGLWGIISASLVMFRSSGLNFLISILGVVIFTGLTAYDVQNI
 SKMDKMLQDDTEIKNRMAVVASLKLDFINLFLYLLRFLGQ
 RRND

t368.aa

TSENQTIKAIIFNSMSFMAMILIQFGLVYAIISGALNKISSNTATALFLLYSALTGVTLSSIFMIYTQGSIVFTFG
 ITAGTFLGMSVYGYTTTDLTKMGSYLI
 MGLWGIISASLVMFRSSGLNFLISILGVVIFTGLTAYDVQNI
 SKMDKMLQDDTEIKNRMAVVASLKLDFINLFLYLLRFLGQ
 RRND

f368.nt

TABLE 1. Nucleotide and Amino Acid Sequences

ATGATCGATTAAACACAAGAAAAACAAGAAATACTAATAAAAACAAGTTTAGCCAAAGTTTCGGGCTTATGT
 CAATTGGACTTTAATCTCAGCAGTATTGCATATGCAACCTCAGAAAATCAAACAATCAAAGCAATAATTCTC
 AAATTCAATGTCATTATGGCTATGATACTTACAAATTGGACTTGTATATGCAATAAGTGGCTCTTAATAAAA
 ATATCAAGCAATACTGCAACAGCTTTCTGCTCTACTCAGCACTAACAGGAGTAACATTATCTTCTATATT
 TGATTTACACACAAGGATCAATAGTATTACACATTGGAATTACTGCTGGAACATTCTTGGAAATGTCTGTTATGG
 ATACACTACAACAACAGATCTAACAAAATGGGAAGCTATTTAATAATGGGTTATGGGAATCATTATTGCATCT
 CTTGTTAATATGTTTTAGAAGCTCAGGTCTTAATTCTTATATCTATTGGCGTAGTTATATTACAGGCT
 TAACAGCTTATGATGTTCAAAATATTCTAAAATGGACAAAATGCTACAAGACGACACTGAAATAAAAACAGAAT
 GCGGTTGTTAGCCTCACTTAACTTTAGATTAAATTATTCTTATATCTTAAGATTGGCCAA
 AGAAGAACGATTAA

t368.nt

ACCTCAGAAAATCAAACAATCAAAGCAATAATTCTCAAATTCAATGTCATTAGGCTATGATACTTACAAAT
 TTGGACTTGTATATGCAATAAGTGGCTCTTAATAAAAATCAAGCAATACTGCAACAGCTCTTCTGCTCTA
 CTCAGCACTAACAGGAGTAACATTATCTCTATATTGATTACACACAAGGATCAATAGTATTACACATTGGA
 ATTACTGCTGGAACATTCTGGAATGTCGTTATGGATACACTAACACAACAGATCTAACAAAATGGGAAGCT
 ATTAAATAATGGGCTTATGGGAATCATTATTGCATCTCTTGTAAATATGTTTTAGAAGCTCAGGTCTTAATT
 CCTTATATCTATTGGCGTAGTTATATTACAGGCTTAACAGCTTATGATGTTCAAAATATTCTAAAATGGAC
 AAAATGCTACAAGACGACACTGAAATAAAAACAGAATGGCGGTTGTTAGCCTCACTTAAACTTTAGATT
 TAAATTATTCTTATATCTTAAAGATTGGCCAAAGAAGAACGATTAA

f371.aa

MKFFFLLQIALILLSNSLLFGQSPPKEKEDSLLYKEGKFKEAILNTLEEIRLNPSNLDARTILWLSIAIGEYK
 RAEKEAIIGLGIKKHDIRIIQALGEAYFFQKNYDNALKYFQEYISLDSKGARIIKVYNLIADSFYELKRYNEADFA
 YEHALRFSPNNQNLLIKLARSRINAKNKLAEELIKILTISPNNLEAKNLLEELKKSNNKP

t371.aa

EDSLLLYKEGKFKEAILNTLEEIRLNPSNLDARTILWLSIAIGEYKRAEKEAIIGLGIKKHDIRIIQALGEAYFF
 QKNYDNALKYFQEYISLDSKGARIIKVYNLIADSFYELKRYNEADFAYEHALRFSPNNQNLLIKLARSRINAKNKL
 LAEEALIKILTISPNNLEAKNLLEELKKSNNKP

f371.nt

ATGAAATTCTTCTATTACAAATAGCTTAATTCTACTATCCAATTCAAGCTTGTATTGGACAATCACC
 GCTAAAGAAAAGAAGACTCTCTCTTCTATATAAAGAAGGAAAATTAAAGAAGCTATTAAACACGTTAGAAGA
 AATTGACTAAATCCTAGTAACCTAGATGCTAGGACAATATTGATATGGAGCTTAATAGCCATAGGAGAAC
 AGAGCTGAAAAGAGGGCATTATAGGACTTGGCATTAAAAACATGACATAAGAATTATTCAAGCACTAGGAGAAC
 CTATTCTTCAAAAAATTATGACAATGCATTAAACTTCAAGAATAACATTAGCCTTGTATTCTAAAGGAGC
 AAAAGAATAATAAAAGTTATAATTAAATTGAGATTCTTTATGAGCTAAAAGATATAATGAAGCCGATTGCA
 TACGAACATGCATTACGTTCTCTCTAATAACCAAAATCTATTAAATAAAATTAGCAAGATCAAGAAC
 AAAATAAAATTAGCAGAAGACACTAATTAAATTCTTACAACTCTCTCTAATAATCTAGAGGCAAAAATT
 ACTAGAAGAATTAAAAAGCAACAAACACCTTGA

t371.nt

GAAGACTCTCTCTTCTATATAAAGAAGGAAAATTAAAGAAGCTATTAAACACGTTAGAAGAAATTGACTAA
 ATCCTAGTAACCTAGATGCTAGGACAATATTGATATGGAGCTTAATAGCCATAGGAGAAC
 AGAGGCATTATAGGACTTGGCATTAAAAACATGACATAAGAATTATTCAAGCACTAGGAGAAC
 CAAAAAAATTATGACAATGCATTAAACTTCAAGAATAACATTAGCCTTGTATTCTAAAGGAGCAAGAAC
 AAGTTTATAATTAAATTGAGATTCTTTATGAGCTAAAAGATATAATGAAGCCGATTGCA
 TACGTTCTCTCTAATAACCAAAATCTATTAAATAAAATTAGCAAGATCAAGAAC
 AAAATAAAATTAGCAGAAGACACTAATTAAATTCTTACAACTCTCTCTAATAATCTAGAGGCAAAAATT
 TTAGCAGAAGACACTAATTAAATTCTTACAACTCTCTCTAATAATCTAGAGGCAAAAATT
 TAAAGCAACAAACACACCTTGA

TABLE 1. Nucleotide and Amino Acid Sequences

f502.aa

MKKANFLSTNFLILLLVCFVNVLFSKDFKFKLVDQFFPFYYKNNKGEYEGLIFSILDWKAKDNNADIMVEHIDN
 LNESEIEDEAIYLGLTYNVKLNDFFYFKSELARSISILFFKNSNKKYKNTHSTFLSNFNFNIGVIKNTIYEDILRLKN
 VNTIFLADNSQELVLALKNDKVDYIYGDKCTLHYIANNFLSEDLVIFTGDFVFSIKNRVAISRNAPEIVKVLNLDL
 FSYLMKMPPEELVFSFLDSNAKGSFVGVLYNDYPPLSFINSQGKLSGILVLDWNLLSRQHIFKPIFKGFSKEDIKK
 SLDGKSVGIFGGIISNDSVLENVYVVKSYDSIFNKNRFLVLAIDNRIYKVIKYILNSIFDDISFESLLQIDKWN
 IVNNNSYGFIEINSITTKYLLKLMGYNGLKSYDSIFNKNRFLVLAIDNRIYKVIKYILNSIFDDISFESLLQIDKWN
 LDKEEINSSRINSYKIMNKVKFNIEEKIWLSKNNKLNLAVKNWYPIDYVEANNYKGINQFLLDKIRMFSLRFNII
 KVHSSLDLKKLIKSGKIDMLNTNATDSNLDNVFNKLNKLNLAVKNWYPIDYVEANNYKGINQFLLDKIRMFSLRFNII
 KSKLILVSSFNEALLLYKGKVDGIISDEYTAAVFEELNIDDVEKIPTFRDLAFDLSLAIYNQDYILKEIIQKVV
 MRSNVDSQMYLNDWKFDIYYKSRSIRFKNFKFLVITFIIFYFTFLGFVIIFMFRLSFEQKRRYSFVMNEKKIAEAA
 NAAKTIFIANVSHDIRTPINGIMAATELLDTTILTVDQKDYVRMINYSSDSLLSLIDDILYLSKIDVNELYVESQE
 IDLESEMEMVLKAFQSQCAKKNIDLFYSKSISIFNNYIKGDIVKIKQVLINLIGNAFKFTDDGVIVLNYEEVCRT
 DGNRVLVTVEFKVIDTGKIEKENFSKIFEIFKQEDDSSSRVHEGAGLGLSISRELIRLMGGLGIAVDSKVGEGETT
 FSFMLPFLLGSELKSKKLSINRFQSVNGDNKVLNVLLSQSKSIKIFEHCSILLGCSSNVRYVASFEDAYKVFKKYPS
 YNFVYINVNNNDNIQEGIRLANNIERLNSDVQIIFLYYLDNKALKNLYGYVKKPLMGLGICSILYKKEFNPEMD
 EDLVPIDSLARIKEPINVLLIAEDNQVNQKVLKDILVVGIGENFIDVVDGVALKSLKDKKYTISFIDIRMPRYD
 GFSVAKEIRKFEKAKNLKPCVLVAVTAHALQEYKDKCLASGMNDYISKPIHISSIKTILKKYLQFEVDDIGENENL
 NQLVKFPNLDVNRALKELNLSYVSYSELCRGLVDFISINIIDLEKAFDEEDLSLIKDISHSISGALSNMRSELYKD
 FQKIEITSKDSISELKKMYSFKDDLFQLISDIKENILFESEIVSENKLYFKNNNDQFLNLLNKLIGIKTRKPREYK
 EILESINKYVLDNNIQVLFSDLRRNRLYRFAESSKILEEIIEMLNKRY

t502.aa

CFVNVLFSKDFKFKLVDQFFPFYYKNNKGEYEGLIFSILDWKAKDNNADIMVEHIDNLNESEIEDEAIYLGLTY
 NVKLNDFFYFKSELARSISILFFKNSNKKYKNTHSTFLSNFNFNIGVIKNTIYEDILRLKNVNTIFLADNSQELVLAL
 KNDKVDYIYGDKCTLHYIANNFLSEDLVIFTGDFVFSIKNRVAISRNAPEIVKVLNLDLFSYLMKMPPEELVFSFLD
 SNAKGSFVGVLYNDYPPLSFINSQGKLSGILVLDWNLLSRQHIFKPIFKGFSKEDIKKSLDGKSVGIFGGIISND
 SVLENVYVVKSYDSIFNKNRFLVLAIDNRIYKVIKYILNSIFDDISFESLLQIDKWNLDKEEINSSRINSYKIM
 NKVKFNIEEKIWLSKNNKLNLAVKNWYPIDYVEANNYKGINQFLLDKIRMFSLRFNIIKVHSSLDLKKLIKSGKI
 DMLNTNATDSNLDNVFNKLNNSRIPLYIFSNKKRVLPSRSLEKFAILDFLYSKNLSNIKSKLILVSSFNEALLL
 YKGKVDGIISDEYTAAVFEELNIDDVEKIPTFRDLAFDLSLAIYNQDYILKEIIQKVVMSNVDSQMYLNDWKFD
 IYYKSRSIRFKNFKFLVITFIIFYFTFLGFVIIFMFRLSFEQKRRYSFVMNEKKIAEAAAKTIFIANVSHDIRT
 PINGIMAATELLDTTILTVDQKDYVRMINYSSDSLLSLIDDILYLSKIDVNELYVESQEIDLESEMEMVLKAFQSQ
 CAKKNIDLFYSKSISIFNNYIKGDIVKIKQVLINLIGNAFKFTDDGVIVLNYEEVCRTTDGNRVLVTVEFKVIDTG
 KGIEKENFSKIFEIFKQEDDSSSRVHEGAGLGLSISRELIRLMGGLGIAVDSKVGEGETTFSFMLPFLLGSELKSK
 LSINRFQSVNGDNKVLNVLLSQSKSIKIFEHCSILLGCSSNVRYVASFEDAYKVFKKYPSYNFVYINVNNNDNIQEGI
 RLANNIERLNSDVQIIFLYYLDNKALKNLYGYVKKPLMGLGICSILYKKEFNPEMDFEDLVPIDSLARIKEPIN
 VLLIAEDNQVNQKVLKDILVVGIGENFIDVVDGVALKSLKDKKYTISFIDIRMPRYDGFSVAKEIRKFEKAKNL
 KPCVLVAVTAHALQEYKDKCLASGMNDYISKPIHISSIKTILKKYLQFEVDDIGENENLQVLFPNLDVNRALKE
 LNLSYVSYSELCRGLVDFISINIIDLEKAFDEEDLSLIKDISHSISGALSNMRSELYKDFQKIEITSKDSISELKKM
 YSFVKDDLFQLISDIKENILFESEIVSENKLYFKNNNDQFLNLLNKLIGIKTRKPREYKEILESINKYVLDNNIQV
 LFSDLRRNRLYRFAESSKILEEIIEMLNKRY

f502.nt

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 TTCAACATTTCATCCAATTTCATATAGGAGTTATAAAAATACAATATGAAAGATATCTAAGGTTAAAAC
 GTTAACACCATTGGCTGATAATTCTCAAGAGTTAGTATTGGCCTTAAAAACGATAAAAGTTGATTATAT

TABLE 1. Nucleotide and Amino Acid Sequences

ATGGTGATTGCAAGACTTACATTATATGCAAATAACTTTAAGTGAAGATCTTGTGATTTTACCGGGATGT
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 TTTGTGGAATCTCTCAAGACAAACATATCTTAAACCTATTAAAGGGATTTCAGGAAATTAAGAAA
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 TAAGATTATAGATTGCTGAGAGCTCAAGATTCTGAAAGAGATTATTGAAATGCTTAATAAGAGATATTA
 G

TABLE 1. Nucleotide and Amino Acid Sequences

TGCTTTGTCAACGTCAATTATTTCTAAGGATATTTCAAGTTAAGCTGTAGATCAATTTCCTTTACT
 ACAAGAATAATAAAGGAGAATATGAAGGACTTATTTCTATTAGATAAAATGGCAAAAGATAATAATGCTGA
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 AATGTAACAAATTAAATGATTTTTATTTAAAGTGAAGCTGCTAGGAGTATTCAATTATTTAAAGACT
 CTAATAAAATATAAAATACCCATTCAACATTATCCATTAAATATAGGAGTTAAAGATAACATATA
 TGAAGATATCTTAAGGTTAAAGCCTAACACCATTTGGCTGATAATTCTCAAGAGTTAGTATTGGCCTTA
 AAAACGATAAAAGTGTGATTATATATGGTATTGCAAGACTTACATTATATTGCAAATAACTTTAAAGTGAAG
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 GGTATCCAATAGATTATGTTGAGGCAAAATAATTAAAGGAATAATCAATTGGTGTGATAAGATTAGAATGTT
 TTCAGGTTGAGATTAAACATAATTAAAGTACACAGCAGTTAGATCTTAAAGTTAAATCAAATCTGGAAAATC
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 GAAAGAAATTATTCAAAGTTGTTATGCGTTCAATGTCAGTCAAGTGTATTAAATGATTGGAAATTGAT
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 GGATGATAATTATTCTCATGTTCTTGTCTTAAATTGATGATATTGATTGTTCTAAAGGTTAGTGTCAA
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 AAATTGGAGCACTGTTGATTGAGCTTATTGGATGCTCTTCAATGTGCGCTATGTAGCGTCTTGTGAGGATGCTTA
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 TATTCTTTGTTAAAGATGATTATTCAACTAAAGCGACATAAGGAAATTAGGTTGAGTGTGAGATTG
 TTAGTGTGAGAACAGCTATTAAATAATGATCAATTAAACCTTCTCAACAAACTTTAATTGGTATTAA

TABLE 1. Nucleotide and Amino Acid Sequences

GACTAGAAAGCCAAGAGAATACAAAGAAATTCTTGAGAGCATTAAATAATGTTTAGACGATAATATTCAGGTA
TTATTTAGTGATCTTCGAGAAATTAAAGATTATAGATTGCTGAGAGCTAAAGATTCTGAAGAGATTATTG
AAATGCTTAATAATAAGAGATATTAG

f527.aa

MNLLVKIAKFILFLFTSCNQKQSEIQLNLTHLLKSSNKNRLDKFLIIDRVVNIYIANKNYEADALEIVNNGIIDDE
SREYYPLYLYLMGNIYDSMGEDFVAFNIVYKRVVDNFDDYVYENHSMKTRVAKKIVNLNIDSIDKINYKFILNMGI
DNLNNEEKGNYFYNLALSLEDVQDYDESYFYYKKFLSIPRAHLKIDSRDYFNVVTKINYFNNPEFVYRNLGDLIQ
DVKNFVLSGNTSKLLNIRDKNNFFIQSWDQKGGKNSINTNSFLTTMIRLGRRKNGIQFAKHLEADSDDISYLE
SRGWDHIEHWYFVFKRIVYPKDPEINNGWTWIGVYLGKK

t527.m

CNQKQSEIQLNLTHLLKSSNKNRLDKFLIIDRVVNIYIANKNYEADALEIVNNGIIDDESREYYPLYLYLMGNIYDSM
GEDFVAFNIVKRVVDNFDDYVYENHSMKTRVAKKIVNLNIDSIDKINYKFILNMGIIDNLNNEEKGNYFYNLALSL
EDVQDYDESYFYYKKFLSIPRAHLKIDSRDYFNVVTKINYFNNPEFVYRNLGDLIQDVKNFVLSGNTSKLLNIRD
KNNFFIQSWDQKGGKNSINTNSFLTTMIRLGRRKNGIQFAKHLEADSDDISYLESRGWDHIEHWYFVFKRIVY
PKDPEINNGWTWIGVYLGKK

f527.nt

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ATAATGGCTGGACTTGGATAGCGTGTATTAGGTAAAAATAA

t527.nt

TGCAACCAAAAGCAAAGCAGATTCAAAATCTTACACATCTTAAATCTCTAATAAAATAGATTAGATAAAT
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TGGAAATTATTGATGATGAATCTAGAGAATATTCTTGTATCTTATTAAATGGCAATATTGATTCATG
GGAGAAGATTGTTAGCTTTAATATTACAAGCGTGTGTTGATAATTGATGATTATGTTATGAAAACCATT
CAATGAAAACAAGGGTTGCTAAAAGATTGTCATTAAATATTGATTCAATCGATAAAATCAATTATTACAAATT
TATATTAAATATGGGAATTGATAATTAAATAATGAGGAAAGGTAATTATTTTATAATCTTGCCTAAGTTG
GAAGATGTTCAAGATTACGATGAATCTTATTGTTATTATAAAAATTCTTCAATTCCAAGGGCACATTAAAAA
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TTTAGGAGATTAACTCAGGATGTTAAAATTGTTCTTCTGGTAATACTCTAAATTGCTTAATATAAGAGAT
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CCACTATGATTAGGCTGGGGGAGAAGAAAAACGGAATACAATTGCAAAGCATCTGAGGCAGATTCTAGTGA
CGATATATCTTATCTTGAGTCAGGGCTGGACCATTATCATGAATGGTATTGTTTAAAGAATTGTTAT
CCTAAAGATCCAGAAATTAAATGGCTGGACTTGGATAGCGTGTATTAGGTAAAAATAA

f541.aa

MNKILLLILLESIVFLSCSGKGS LGSEIPKVS LIIDGT FDDKS FNE SALNGVKKVKEEFKIELVLKESSNSY LSD
LEG LKDAGSDLIW LIGYRF SDVAKVA ALQNPDMKYAIIDPI YSN DPIP ANLVGMTFRAQEGAFLTGYIAAKLSK TG

TABLE 1. Nucleotide and Amino Acid Sequences

KIGFLGGIEGEIVDAFRYGYEAGAKYANKDIKISTQYIGSFADLEAGRSVATRMSDEIDIIHHAAGLGGIGAIEV
 AKELGSGHYIIGVDEDQAYLAPDNVITSTTKDVGRLNIFTSNHLKTNTFEGGKLINYGLKEGVVGFVRNPKMISF
 ELEKEIDNLSSKIINKEIIVPSNKESEYKFLKEFI

t541.aa

CSKGKSLGSEIPKVSLIIDGTFDDKSFNEALSNGVKKVKEEFKIELVLKESSNSYLSDEGLKDAGSDLIWLIGY
 RFSDVAKVAALQNPDMKYAIIDPIYSNDPIPANLVMGTFRQEGAFLTGYIAAKLSKTGKIGFLGGIEGEIVDAFR
 YGYEAGAKYANKDIKISTQYIGSFADLEAGRSVATRMSDEIDIIHHAAGLGGIGAIEVAKELGSGHYIIGVDEDQ
 AYLAPDNVITSTTKDVGRLNIFTSNHLKTNTFEGGKLINYGLKEGVVGFVRNPKMISFELEKEIDNLSSKIINKE
 IIVPSNKESEYKFLKEFI

f541.nt

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t541.nt

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 ACCTGAAAGCTGGTAGAGCGTTGCAACTAGGATGTAATTCTGATGAGATAGACATTATTCTACATGCTGCAGGCC
 TGGAGGAATTGGGCTATTGAGGTTGCAAAGAACTGGTCTGGCATTACATTATTGGAGTTGATGAAGATCAA
 GCATATCTGCTCCTGACAATGTAATAACATCTACAACTAAAGATGTTGGTAGAGCTTAAATATTTCATCTA
 ACCATTAAAACATAACTTCTGAAGGTGGCAAATTAAATAATTATGGCCTTAAAGAAGGAGTTGTGGGTTGT
 AAGAAATCTAAAATGATTCTTGAACCTGAAAAGAAATTGACAATCTTCTAGCAAAATAATCAACAAAGAA
 ATTATTGTTCCATCTAATAAAAGAAAGTTATGAGAAGTTCTTAAAGAATTATTAA

f561.aa

MYKNGFFKNYLSLFLIFLVIACTSKDSSNEYVEEQAENSSKPDDESKIDEHTIGHVFHAMGVVHSKKDRKSLGKNI
 KVFYFSEEDGHFQTIPSKENAKLIVYFYDNVYAGEAPISISGKEAFIFVGITPDFKKIINSNLHGAKSDLIGTFKD
 LNIKNSKLEITVDENNSDAKTFLESVNYIIDGVEKISPMLTN

t561.aa

TABLE 1. Nucleotide and Amino Acid Sequences

CTSKDSSNEYVEEQEAEENSSKPDDSKIDEHTIGHVFHAMGVVHSKKDRKSLGKNIKFVYFSEEDGHFQTIPSKEA
KLIVYFYDNVYAGEAPISISGKEAFIFVGITPDFKKIINSNLHGAKSDLIGTFKDLNIKNSKLEITVDENNSDAKT
FLESVNYIIDGVEKISPMLTN

f561.nt

ATGTATAAAAATGGTTTTTAAACTATTGTCATTGTTTAATTAGTTAGTAATTGCTGTACTCAAAAG
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TACTATTGGCACGTTTCACGCTATGGGAGTAGTTCAATTCAAAAAGGATCGAAAAGTTGGGAAAAATATA
AAGGTTTTTATTTCTGAAGAAGATGGACATTCAAAACAATACCCCTCAAAAGAGAATGCAAAGTTAATAGTT
ATTTTATGACAATGTTATGCAGGAGAGGCTCAATTAGTATCTCTGGAAAAGAACGCTTATTGGTACTTTAAAGAT
TACCCCTGACTTAAAGATTATAAATAGCAATTACATGGCGCTAAAGTGTATCTGGTACTTTAAAGAT
CTTAATATTTAAAGATTCAAAATTGAAATTACAGTTGATGAGAATAATTCAAGATGCCAAGACCTCTGAATCTG
TTAATTACATTATCGACGGCGTTGAAAAAATTACCTATGTTAACGAATTAA

t561.nt

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GGGAAAAAATATAAAGGTTTTTATTTCTGAAGAAGATGGACATTCAAAACAATACCCCTCAAAAGAGAATGCA
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TTTTGTTGGATTACCCCTGACTTAAAGATTATAAATAGCAATTACATGGCGCTAAAGTGTATCTTATTGG
TACTTTAAAGATCTAATATTAAAATTCAAAATTGAAATTACAGTTGATGAGAATAATTCAAGATGCCAAGACC
TTCCTGAATCTGTTAATTACATTATCGACGGCGTTGAAAAAATTACCTATGTTAACGAATTAA

f604.aa

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KPIKNAKHN

t604.aa

CNNNSEKEKLAFLKVVYIGGAPSSLDPLVDETIGARILEQIFSGLLTLNTGKLKPGLAKNWEASKDKKTYQFYLR
DNLFWSDGVEITAEGIRKSFLRILNKETGSTNVDMILKSIKNGQEYFDGKVSDSELGIKAIDSKTLTAPKPY
FLELLLHYAFMPVPIHVIEKYGNWTSPEMVTSGPFLKKRLPNEKIIFEKNERYYNAKEVELDELVYITSNDL
TVYNMYKNNEIDAIFNSIPPDIVNEIKLQKDYYQHKSNAIYLYSFNTKIKPLDDARVREALTLAIDRETLTYKVLN
DGTVPTRITPDLKNYNYGKKLALFDPEKSKLLADAGYPNGKGPMLTLYKNTNETHKKIAAFIQNQWKKILNIN
LMLTNENWPVLTNSRNTGNFEIRVGRIGEYLDPTHYFTIFTRENSQLASYGYSNLEFDKLIRESDEKDPPIKRQ
LLRKAESIIIEKDFPAAPIYIYSGHYLFRNDKWTGWNPNVSEVYIYSELKPIKNAKHN

f604.nt

ATGAGCTTTAATAAAACTAAAAAAATCGGTTAAAGAAATTAAAGTAAACACTACTTATGCTTGCTGTCTTAA
TTGCATGCAATAATAATTCAAGAAAAAGAAAAATTAGCATTTAAAGTATACATAGGGGGAGGCCCTCATCGCTTGA
CCCTCATTGGTAGATGAGACAATAGGAGCAAGAATTAGAACAAATTCTCAGGGCTTTGACATTAAATACC
AAAACAGGAAAGCTAAAGCCCGACTGCTAAAATGGGAAGCCTCAAAAGATAAAAAACATATCAATTATC
TAAGGGACAACCTTTTGGAGCGATGGAGTTGAAATTACCGCTGAAGGGATAAGAAAATCTTTTAAGAATT
AAATAAAGAAACAGGATCTACAAATGTTGACATGCTAAATCAATAAAAAATGGACAAGAGTATTTGACGGG
AAAGTATCCGATTCTGAACCTGGAAATCAAGGCAATTGATAGTAAACGCTGGAAATAACACTTACGGCCCCAAAGC
CATATTCTTGAACCTGCTTACATTACCGATTACGCCAGTACCTATTGATTGAAAAATATAAGGGAAA

TABLE 1. Nucleotide and Amino Acid Sequences

TTGGACAAGCCCTGAAAACATGGTTACTAGCGGTCTTTAAATTAAAAAAAGATTACCTAATGAAAAAATTATC
 TTTGAAAAAAACGAACGTTATTATAATGCAAAAGAAGTAGAACTTGATGAGCTTGCTACATTACGTCTGACAATG
 ATCTTACTGTGTACAATATGTACAAAACAGAAATTGATGCTATTTAACAGCATCCGCCGACATTGTA
 TGAAATAAAACTACAAAAGACTATTACCAACACAAAAGTAATGCAATTATTTATATTCAATTAAACAAAATA
 AAACCCCTGATGATGCTAGAGTTAACCTAGCTATTGACAGAGAAACTTTAACTACAAAGTGC
 TAAATGATGGCACAGTCCCTACAAGAGAAATACTCCTGATCTAAAATTACAATTACGGTAAAAATTGGCTT
 ATTTGATCCTGAAAATCTAAAAGCTTGGCAGATGCAGGGTATCCTAATGGAAAGGATTCCAATGCTAAC
 CTAAAATATAATACAAACGAAACTCATAAAAATTGCTGATTTATTCAAAACCAATGGAAAAAAATTCTAAATA
 TCAATCTTATGCTTACCAACGAAAATTGCCCTGTTTACCAACAGCAGAAATACTGGCAATTGAAATAAAAG
 AGTTGGACGCATTGGGAATTAGATCCACACACATACTTACTATTCACAAGAGAAAATTACAACATTGCA
 TCATACGGATATTCAACCTAGAATTGACAAACTCATCAGAGAATCAGATCTGAAAAGATCCTATAAAAGAA
 ACAATTACTCAGAAAAGCAGAATCAATAATTGAAAAGATTTCCTGCTGCACCAATATACATATATTCTGG
 GCATTATCTTTAGAAACGATAATGGACTGGATGGAATCCTAATGTATCAGAGGTTATTATCTTCTGAATTA
 AAACCAATTAAAATGCAAAACATAATTAA

t604.nt

TCGAATAATAATTCAAGAAAAAGAAAATTAGCATTAAAGTATACATAGGGGAGGCCCTCATCGCTTGACCTC
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 TAAACTACAAAAGACTATTACCAACACAAAAGTAAATGCAATTATTTATATTCAATTACAAAATAAAACC
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 GATGGCACAGTCCCTACAAGAGAAATACTCCTGATCTTAAACATTACGGTAAAAATTGGCTTATTG
 ATCCTGAAAATCTAAAAGCTTGGCAGATGCAGGGTATCCTAATGGAAAGGATTCCAATGCTAACACTAAA
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 CGGATATTCAAACCTAGAATTGACAAACTCATCAGAGAATCAGATCTGAAAAGATCCTATAAAAGAAAACAA
 TTACTCAGAAAAGCAGAATCAATAATTGAAAAGATTTCCTGCTGCACCAATATACATATATTCTGGCATT
 ATCTTTTAGAAACGATAATGGACTGGATGGAATCCTAATGTATCAGAGGTTATTATCTTCTGAATTAAC
 AATTAAAATGCAAAACATAATTAA

f736.aa

MKKVIIILIFMLSTSLLYNCKNQDNEKIVSIGGTTVSPILDEMILRYNKINNNKVTYDAQGSSVINGLFNKIYK
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 SYSSIKDLLLNLKIFKTHEEAQFRQDGIVVKSNGEVIEKTSLTPHSIGYIGLGYAKNSIEKGLNILSVNSTYPTKET
 INSNKYTIKRNLIIVTNNKYEDKSVTQFIDFMTSSTGQDIVEEQGFLGIKT

t736.aa

CKNQDNEKIVSIGGTTVSPILDEMILRYNKINNNKVTYDAQGSSVINGLFNKIYKIAISSRDLTKEEIEQGAK
 ETVFAYDALIFITSPEIKITNITEENLAKILNGEIQNWKQVGGPAKINFINRDSSSGYSSIKDLLLNLKIFKTHE
 EAQFRQDGIVVKSNGEVIEKTSLTPHSIGYIGLGYAKNSIEKGLNILSVNSTYPTKETINSNKYTIKRNLIIVTNN
 KYEDKSVTQFIDFMTSSTGQDIVEEQGFLGIKT

f736.nt

TABLE 1. Nucleotide and Amino Acid Sequences

ATGAAAAAAGTTATTATCTTAATTTTATGCTATCAACAAGTTATTATACAACGTAAAAATCAAGACAATGAAA
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 CAATAACTAAAGTAAACATACGATGCACAAGGAAGTAGTGTGGCATAAACGGCTATTAAACAAAATATAAA
 ATAGCAATATCATCAAGAGATTAAACAAAAGAAGAATTGAACAAGGGGAAAAGAAAATCTGTATTTGCTTATGATG
 CTTAATTTCACTTACAAGGCCGAAATAAAATTACAAATATTACAGAAGAAAATCTAGCTAAAATACTAAATGG
 AGAAAATTCAAAATTGAAACAAGTGGGAGGTCCTGATGCTAAAATCAACATTCAATCGAGACTCTCTCTGGT
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 CTCATTATTGATTGATTCATGACAAGCTCAACTGGACAAGATATTGTTGAAGAACAAAGGTTTTAGGGATAAAAAC
 ATAA

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TGTAAAATCAAGACAATGAAAAAATTGTATCAATTGGAGGATCTACAACTGTAAGCCAATACTAGACGAAATGA
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 GCTATTAAACAAATATAAAATAGCAATATCATCAAGAGATTAAACAAAAGAAGAATTGAACAAGGGGAAA
 GAAACTGTATTGCTTATGATGCTTAATTTCATTACAAGCCCTGAAATAAAAATTACAAATATTACAGAAGAAA
 ATCTAGCTAAAATACTAAATGGAGAAATTCAAAATTGAAACAAGTGGGAGGTCCTGATGCTAAAATCAACTTTAT
 CAATCGAGACTCTCTGTTCTTATTCGTCTATAAAAGACCTACTTCTTAATAAAATATTCAAAACTCACGAA
 GAAGCTCAATTAGACAAGACGGAATAGTGGTAAATCTAATGGAGAGGTAAATTGAAAAACAAGCCTACTCCCC
 ACTCAATAGGATATATAGGTCTGGATACGAAAAATTCAATAGAAAAGGGTTGAATATTCTTCTGTTAACAG
 CACATATCCTACAAAAGAAACAATAATAGCAATAACACCATTAAAAGAAATTAAATAGTTACAATAAC
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 AAGGTTTTAGGGATAAAAACATAA

f752.aa

MNKKLNEVLLKLDQDLIKCVKGSLDLEISGVTYSSKLVLPRFVFFALPGIHFDFHDFIEIAIQGSNVVVCSDVD
 FYSPNVTYIKVDDFNIRKFMNSNFSNIFYDEPSKKLKVGVTGTDGKSSVCYYIYLLFKKKGVKVGFI
 GSTVFFDDGSLSIKNPYRQSTPESTEIHSFLSTMVKNEAQYIALESTSHGLDLETARLIDVNYFAVVFTNIGHELEFHGT
 IONYLNVKLGLFRSVSDAGFG
 LNVKLGLFRSVSDAGFGVINLDDLYSSDFKNAVKKSFYSLKSSKADFFVFSIDEKTDSTRF
 EFYHKGVKYLANVSLGSFNVENVMALILV
 SLGSFNVENVMALILVSQLIQLNIDIQDIVDKLNCIKSLDGRMDSINLGQNF
 SVIIDYHTPGAFSKLFP
 IFKRFA
 TNRLISVFGSAGERDVEKRF
 LQGQIADIYSDLII
 LCDEDPRGENSMCI
 IKDI
 AKGIVNKVENKDLFFI
 IADRQ
 KQIAIE
 KAISLAKAGDLVVALGKG
 HESSIIYKNREVFWNEQEVVKNAI
 LSLEKSEKEK

t752.aa

CVKGSLDLEISGVTYSSKLVLPRFVFFALPGIHFDFHDFIEIAIQGSNVVVCSDVD
 FYSPNVTYIKVDDFNIRK
 FMSNFSNIFYDEPSKKLKVGVTGTDGKSSVCYYIYLLFKKKGVKVGFI
 GSTVFFDDGSLSIKNPYRQSTPESTEI
 HSFLSTMVKNEAQYIALESTSHGLDLETARLIDVNYFAVVFTNIGHELEFHGT
 IONYLNVKLGLFRSVSDAGFG
 VINLDDLYSSDFKNAVKKSFYSLKSSKADFFVFSIDEKTDSTRF
 EFYHKGVKYLANVSLGSFNVENVMALILV
 SQLIQLNIDIQDIVDKLNCIKSLDGRMDSINLGQNF
 SVIIDYHTPGAFSKLFP
 IFKRFA
 TNRLISVFGSAGERDVEK
 RFLQGQIADIYSDLII
 LCDEDPRGENSMCI
 IKDI
 AKGIVNKVENKDLFFI
 IADRQ
 KQIAIE
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 HESSIIYKNREVFWNEQEVVKNAI
 LSLEKSEKEK

f752.nt

ATGAATAAAAATTAATGAAGTTTATTAAAGTTAGATCAAGATTAAATAAAATGTGAAAAGGTTCTTGATT
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 TTTTTTATGATGAGCCTCAAAAAAATTAAAAGTTATTGGAGTCAGTGGCACTGACGGAAAAGTTCTGTTGTTA
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 GGAAGCTTGTATTAAAATCCTACAGACAATCAACTCCGAGTCTACGGAAATACATTCAATTAAAGCACCATGG

TABLE 1. Nucleotide and Amino Acid Sequences

TTAAAAATGAAGCTCAATATGCAATTCTGAATCTACTTCTCATGGGCTTGACCTGAAACAGCAAGGCTTATTGA
 TGTTAATTATTTGCAGTTGTTTACCAATATTGGACATGAGCATTGAAATTCTATGGCACATTCAAAATTAT
 TTGAATGTCAAGCTGGGTCTTTTCGGTCTGTTAGTGTGATGCTGGTTGGGTATTAAATCTTGTGACCTTT
 ATTCTTCTGATTTAAGAATGCTGTTAAGAAATCTTACTTATAGCTTAAAAGCAGTAAGCGGATTTTTGT
 TAGTTTATTGATGAGAAAACGATTCTACTAGATTGAATTATCACAAAGGGGTTAAATATCTTGTAAATGTT
 AGCCTACTGGGAGTTAATGTTGAGAATGTAATGGCTCTTATTTAGTTCTCAAATTAAATATCGATA
 TTCAAGATATTGTTGATAAACTTAAGTGCATTAAAAGTCTTGATGGCGTATGGATAGTATTAAATTGGGCAAA
 TTTTCTGTAAATAATTGATTATGCTCATACTCCTGGTCTTCCAAGCTTTCTATTAAAGATTTGCT
 ACCAATAGATTGATTCTGTTTGGCTCTGCAGGAGAAAGAGATGTTGAAAAAAAGATTTGCAAGGGCAAATCG
 CAGATATTATTCTGATTTAATAACTTGCATGAAAGATCCAAGAGGAGAAATAGTATGTTATAATTAAAGA
 CATTGCAAAGGAATTGAAATAAAAGTGAAGAAATAAGGATTATTATTGCTGATAGAAAGCAGGCTATTGAA
 AAAGCAATAAGTCTGCAAAGCAGGAGATTGGTTGTTGCTTGGCAAAGGTCAAGGTTCAATAATTATA
 AAAATAGAGAAGTTTTGGAATGAACAAGAGGTAGTTAAATGCTATTAAAGTTAGAAAAATCAGAAAAGGA
 GAAGTGA

t752.nt

TGTGAAAAGTTCTTGATTTAGAAATATCAGGAGTTACTTATAGTTCAAATTGGTTTGCCAGGTTGT
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 TGTGTTCACGAGATGTGGATTTACAGTCTAAATGTTACTTATATTAAAGGTAGATGACTTAAACATAAGAAA
 TTATGTCTAATTTCAAATATTGATGAGCCTTCAAAAAAAATTAAAGTTATTGGAGTCAGTGGCACTG
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 GTTAGAAAATCAGAAAAGGAGAAGTGA

f798.aa

MVFRYKHLELIMLPMLMSCAFFKKPQSVHQDSNTGKPIISDEKLHLISGKISNKKLPIINSNHDVTWIKTKAMTI
 LGEDGKEIPEFKNKFGYSYIISPVKMDGKYSYYASLLILFETTKNGDDEYEIEVDVKFVTAGSTLELKNSLLAVENS
 QEEGYVTAYPFGILMSDEIKNAFKLTYKNHWNYMLADLTVKNKLQETKIKYKISLNSKLIIEFLKEVLKENSILK
 DIAGDLFEDI

t798.aa

CAFFKKPQSVHQDSNTGKPIISDEKLHLISGKISNKKLPIINSNHDVTWIKTKAMTI
 LGEDGKEIPEFKNKFGYSYIISPVKMDGKYSYYASLLILFETTKNGDDEYEIEVDVKFVTAGSTLELKNSLLAVENS
 QEEGYVTAYPFGILMSDEIKNAFKLTYKNHWNYMLADLTVKNKLQETKIKYKISLNSKLIIEFLKEVLKENSILK
 DIAGDLFEDI

f798.nt

ATGGTATTAGAACATATAAACATTGGAACTAATAATGCTGCCATGTTAATGCTGAGTTGCGCTTTTTAAGA
 AACACAACTCTGTACATCAAGACAGCAAACTGGCAAACCAATAAGCGATGAAAATACATTAAATATCAGGCAA
 AATTCAAATAAAAATTGCCAATCATAAATAGTAATCATGACGTAATTGGATAAAAACAAAGGCAAATGACAATC

TABLE 1. Nucleotide and Amino Acid Sequences

TTAGGCAGAGATGGAAAAGAAATACCAAGAATTAAAAACAAATTGGATATTCTTATATAATATCCTGTAAAAA
 TGGATGGAAAATATAGTTATTACCGTCATTATTAATACCTTTGAAACAACTAAAATGGAGATGATGAATATGA
 AATTGAAGATGTTAATTGTAACAGCTGGTCCACCCCTAGAACTAAAAATTCTCTTAGCTGTTGAAAATTCA
 CAAGAAGAAGGATATGTTACTGCATACCCATTGGAATTGATGAGTGACGAGATTTAAATGCTTTAAATTAA
 CATATAAAAATGGTCATTGGAATTATGCTTGAGATTTAACTGTCAAAAATAACTCAAGAAACTAAAAT
 TTATAAAAATTCTCTTAATTCAAAATTAAATTATTGAATTTTAAAAGAAGTGCTAAAAGAAAATTCTATATTAAAA
 GACATAGCTGGAGATTGAAAGATATATAA

t798.nt

TGCGCTTTTTAAGAAACCACAATCTGTACATCAAGACAGCAATACTGGCAAACCAATAAGCGATGAAAATTAC
 ATTTAATATCAGGAAAATTCAAATAAAAATTGCAATCATAAATAGTAATCATGACGTAACCTGGATAAAAC
 AAAGGCAATGACAATCTTAGGCGAAGATGGAAAAGAAATACCAAGAATTAAAAACAAATTGGATATTCTTATATA
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 AGCTGTTGAAAATTCAACAAGAAGGATATGTTACTGCATACCCATTGGAATTGATGAGTGACGAGATTA
 AATGCTTTAAATTAAACATATAAAAATGGTCATTGAAATTATGCTTGAGATTTAACTGTCAAAAATAACTTA
 CTCAAGAAACTAAAATTATAAAATTCTCTTAATTCAAAATTAAATTATTGAATTTTAAAAGAAGTGCTAAAAGA
 AAATTCTATATTAAAAGACATAGCTGGAGATTGAAAGATATATAA

f805.aa

MLRKLKDISKIVLVTDGLTPNCQTCGKLIANGDEVYIAEDGLFHSVKSNTIAGSTLTMIQGLKNLIEFGFSLSDAV
 QASSYNPTRILNIDKKGLICHGYDANLVLDKDFNLKLTMIESKIIIFNNL

t805.aa

CQTCGKLIANGDEVYIAEDGLFHSVKSNTIAGSTLTMIQGLKNLIEFGFSLSDAVQASSYNPTRILNIDKKGLICH
 GYDANLVLDKDFNLKLTMIESKIIIFNNL

f805.nt

ATGCTTAGAAAGCTAAAGATAAGTAAAGTACTGACGGACTTACTCCGAATTGTCAAACTTG
 GAAAACAAATTGCAAACGGAGACGAAGTTATATTGAGAAGATGGATTATTCCATAGCGTAAAAGCAACACAA
 ATAGCTGGATCAACACTCACAATGATAAGGTCTTAAATTAAAGAATTGGTTCAGCTTAAGCGATGCTGTT
 CAAGCAAGCTCTTACAATCCAACAAGAATTCTCAATATTGATAAAAAGGGCTTAATATGTCATGGATATGTC
 AACCTCAATGTCCTAGATAAAGATTAACTAAAGTTAACATGATAGAATCTAAAATAATTAAACAATCTCTA
 A

t805.nt

TGTCAAACCTGTGGAAAACCTAATTGCAAACGGAGACGAAGTTATATTGAGAAGATGGATTATTCCATAGCGTGA
 AAAGCAACACAATTGCTGGATCAACACTCACAATGATAAGGTCTTAAATTAAAGAATTGGTTCAGCTT
 AAGCGATGCTGTTCAAGCAAGCTCTTACAATCCAACAAGAATTCTCAATATTGATAAAAAGGGCTTAATATGTC
 ATGGATATGTCAAACCTCAATGTCCTAGATAAAGATTAACTAAAGTTAACATGATAGAATCTAAAATAATT
 TTAACAATCTCTAA

f635.aa

MKILWLIILVNLFLSCGNESKEKSNLGLRLRELEISGGGSESKIEVYKEFIEKEDKNILKIVNSIDKKARFFNLIG
 LEFFKLGQYGPAYEYFAKNLEINPNNYLHFYIGVASYNLAKNLRVKDEVEKYIILAEFLKSLSI
 RDDFKDSLFAISNMVYDLDKQLEAKNYLNKLGDGEDYFEFLMLRGANYYSLGDLGNAILFYDKASKKASTEEQKEGVSRIMSN
 LK

t635.aa

TABLE 1. Nucleotide and Amino Acid Sequences

CGNESKEKSNLGLRLRELEISGGSESKEVYKEFIEKEDKNILKIVNSIDKKARFFNLIGLEFFKLGQYGPAlEY
 FAKNLEINPNNYLSHFYIGVASYNLAKNLRVDEVEKYIILAENSFLKSLSI RDIFKDSLFAISNMYVYDLDKQLE
 AKNYLNKLGDMGEDYFEFLMLRGANYYSLGDLGNAILFYDKASKKASTEEQKEGVSRIMSNLK

f635.nt

ATGAAAATTTGTGGTTAATAATTCTTGTAAATTATTTATCTTGTGGCAATGAATCTAAAGAAAATCAAATC
 TTGGTCTTAGATTAAGAGAATTGGAATTTCAGGTGGATCTGAATCTAAGATTGAAGTTATAAGAATTAT
 TGAAAAGAAGATAAGAATTAAAGATAGTTAATTCCATTGATAAGAAAGCCAGATTTTAATTAAATTGGT
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 ATTATTTATCTCATTAAATAGGTGTTCTTATAATTAGCTAAAGAGTAAAGATGAAGTTGA
 AAAATACATAATTCTGCTGAAATTCTTTAAATCACTTCATTAGAGATGATTAAAGATTCTCTTTT
 GCCATTCTAATATGTACGTATGATCTGATAAACAACTTGAAGCTAAAATTATTAAATAACTTGGTGATA
 TGGTGAGGACTATTGTGAGTTAAATGTTAAGAGGTGCAAATTATTTCGCTGGCGATCTGGTAATGCTAT
 ATTGTTTATGATAAAGCTAGTAAAAGGCTCAACTGAAGAGCAAAAGAAGGTGTTCTAGGATCATGAGTAAT
 TTGAAGTAA

t635.nt

TGTGGCAATGAATCTAAAGAAAATCAAATCTGGTCTTAGATTAAGAGAATTGGAATTTCAGGTGGATCTG
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 CTAAAATTAAAGAGTAAAGATGAAGTTGAAAATACATAATTCTGCTGAAATTCTTTAAATCACTTC
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 GCTAAAATTATTAAATAAACTTGGTATATGGTGAGGACTATTGAGTTAAATGTTAAGAGGTGCAAATT
 ATTATCGCTGGCGATCTGGTAATGCTATATTGTTATGATAAAGCTAGTAAAAGGCTCAACTGAAGAGCA
 AAAAGAAGGTGTTCTAGGATCATGAGTAATTGAGTAA

f314.aa

MNNCLIKFFIFLLVFSNSYVAFSKNVNLIVTAMDSEFDQINKLMSNKEEIVLKEYGLNKKILKGKLSNRNVMI
 CGVGKVNAGWTSYILSKYNISHVINSGVAGGVVSAYKDIKVGDVVSSEVAYHDVDLTKFGYKVGQLTGLPQK
 FNANKNLIKNAIEAIKSKVGGSNAYSGLIVSGDQFIDPTYINKIIGNFKDVIAMEGAAIGHVSHMFNIPFIVIR
 SISDIVNKEGNEVEYSKFSKIAAFNSAKVQEIILRKLZ

t314.aa

KNVNLIVTAMDSEFDQINKLMSNKEEIVLKEYGLNKKILKGKLSNRNVMIICGVGKVNAGWTSYILSKYNISH
 VINSGVAGGVSAKYKDIKVGDVVSSEVAYHDVDLTKFGYKVGQLTGLPQKFNANKNLIKNAIEAIKSKVGGSN
 AYSGLIVSGDQFIDPTYINKIIGNFKDVIAMEGAAIGHVSHMFNIPFIVIRSIISDIVNKEGNEVEYSKFSKIAA
 FNSAKVQEIILRKLZ

f314.nt

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 TGTGGGTTGTAAGTTAATGCTGGTGTGGACTAGCTACATTGCTAAATACAAACATAAGTCATGTCATTA
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 CAGCCAAAGTTGACAAGAAATTAAAGAAAATTAA

TABLE 1. Nucleotide and Amino Acid Sequences

t314.nt

AAAAATGTCATGTTAATAGTAACGTGCTATGGACTCTGAGTTGATCAGATAAATAAGCTATGTCTAATAAGG
 AAGAAATAGTCTTAAGGAGTATGGTCTTAATAAAAAGATTTAAAGGGGAAGTGTCTAATCGCAATGTATGGT
 TATTATTTGTGGGTTGGTAAGGTTAATGCTGGTGTGGACTAGCTACATTTGTCAAAATACAACATAAGTCAT
 GTCATTAATTCTGGCGTTGCTGGCGTTGTTAGTGCATAACAAAGATATTAAAGTGGGAGATGTGGTGTG
 CTTCAGAGGTTGCATATCATGATGTTGACTAAATTGAGATAACAGGTAGGACAGCTTACAGGAGGATTGCC
 TCAAAAATTAAATGCCAATAAAAATTAAATAAGAATGCCATAGAGGCCATTAATCAAAGGTTGGAGGTTCTAAT
 GCATATTCAAGGATTAATAGTTCAAGGAGATCAGTTATTGATCCAACCTTATTTAAACAATTATAGGAAACTTTA
 AAGATGTAATAGCTGTTGAGATGGAAGGTGCAGCAATAGGGCATGTTCTCATATGTTAATATACCTTTATAGT
 TATTAGGTCAATATCTGACATTGAAATAAAGAAGGGATGAGGTTGAATATAGTAAATTCTAAAATAGCTGCT
 TTCAATTCAAGCCAAAGTTGTACAAGAAATTAAAGAAAACCTTAA

f32.aa

MNTKTLYLISLILLACNKNKIPLIQKLDLPKSSILGFSNKMGIKKDYAFLSKSTKNSELDYDYAILLRKDEVV
 KIEKLEKTERYGIEGNWLVNYKGTKRYIFSKDINIVNNLIIIDHSKZ

t32.aa

CNKNKIPLIQKLDLPKSSILGFSNKMGIKKDYAFLSKSTKNSELDYDYAILLRKDEVVKIEKLEKTERYGIE
 GNWLVNYKGTKRYIFSKDINIVNNLIIIDHSKZ

f32.nt

ATGAATACAAAACATTATTTAATATCCTTAATTCTTTAGCTTGCATAAAAAATAACAAAATTCCCTCTCATT
 AAAAATTAGATTGCCAAAAGCAGCATTCTGGCTTAGCAATAAAATGGGCATAATAATAAAAGATTATGCTTT
 TCTTAGTAAAGCACTAAGAAAATAGCAATTGGATTATGATACGCATTCTACTCAGAAAAGACGAAGTCGTA
 AAAATTGAAAAAACACTAGAAAAACAGAGCGCTATGGAATTGAGGAAATTGGATCTAGTCAATTACAAGGAA
 CTAAAAGATACATCTTAGCAAAAGACATCAATATAGTCAACAAATTAAATTGATCATTCTAAATAG

t32.nt

TGCAATAAAAATAACAAAATTCTCTCATTCATAAAATTAGATTGCCAAAAGCAGCATTCTGGCTTAGCAATA
 AAATGGGCATAATAATAAAAGATTATGCTTTCTTAGTAAAGCACTAAGAAAATAGCAATTGGATTATGATTA
 CGCAATTCTACTCAGAAAAGACGAAGTCGAAAAATTGAAAAACACTAGAAAAACAGAGCGCTATGGAATTGAA
 GGAAATTGGATCCTAGTCAATTACAAGGAACTAAAAGATACATCTTAGCAAAAGACATCAATATAGTCAACAAATT
 TAATAATTGATCATTCTAAATAG

f320.aa

MKSIYALLFLFINLSLLANNISKDKLEVLLKIAQAMNKECKNFI EKNPIQFLKEIKPLVDAEKNNLTLINKKIP
 PENYKIPDLVNIDDFEDLKNLGAKTIKVRKILIEDLIRLIKDAKKFGIEIKIKSAYRTQEQYQKFLFDYNVKT
 YGRKVAETQSAIPGHSQHHMGT AIDFINIDDNLLNTKEGKWL
 YENSLKYGFSVSYPKGYETDTGYKAEPWHYLYIGPKPCFIQKKYFNNLQHKLLEFW
 NQNKTNLINLIEKYANZ

t320.aa

NNISKDKLEVLLKIAQAMNKECKNFI EKNPIQFLKEIKPLVDAEKNNLTLINKKIP
 PENYKIPDLVNIDDFEDLKNLGAKTIKVRKILIEDLIRLIKDAKKFGIEIKIKSAYRTQEQYQKFLFDYNVKT
 YGRKVAETQSAIPGHSQHHMGT AIDFINIDDNLLNTKEGKWL
 YENSLKYGFSVSYPKGYETDTGYKAEPWHYLYIGPKPCFIQKKYFNNLQHKLLEFW
 NQNKTNLINLIEKYANZ

f320.nt

TABLE 1. Nucleotide and Amino Acid Sequences

ATGAAATCAATTATGCTTATTATTCTATTAAATTATCTTTGTGGCTAACAAACATTCAAAAAAAGATT
 TAGAAGTACTGCTAAAGATTGCCAAGCAATGAATAAGGAATGCAAAATTTATTGAAAAAAATCCTATTCAAGTT
 CTTAAAAGAAATAAAACCTTAGTAGATGCAGAAAAAAATAACCTCTTAACCTAAATAAAATAACCAATT
 CCTGAAAATTATAAAATACCTGATCTGGTAAATTGATGATTTGAAGATCTTAAATCTTGGAGCAAAGACTA
 TTAAAGTAAGAAAATATTAATCGAAGATTAACTCGACTAATAAAAGATGCAAAAAAATTGGGATTGAAATTAA
 AATCAAATCTGCTTACAGAACGCAAGAATATCAAATTTTATTGATTACAATGTCAAAACATTATGGCAGAAAA
 GTTGCAGAAACCCAATCAGCAATTCCAGGCCATTCTCAACATCATGGAACAGCAATAGATTATAAATATAG
 ATGATAATTACTAAACACAAAAGAAGGAAATGGCTTATGAAAACCTCTTAAATACGGATTTCCGTTCTATA
 CCCAAAAGGATATGAAACGGACACTGGATATAAGCAGAGCCTGGCACTACTTATACATAGGACCTAACCCATGC
 TTTATTCAAGAAAATATTTAATAATTACAACATAAGCTTCTGAATTGGAACAGAACAAAACAAATCTTA
 TTAACCTAATTGAAAAATATGCAAACCTAA

t320.nt

AACAACATTCAAAAAAAGATTAGAAGTACTGCTAAAGATTGCCAAGCAATGAATAAGGAATGCAAAATTTA
 TTGAAAAAAATCCTATTCACTTAAAGAAATAAAACCTTAGTAGATGCAGAAAAAAATAACCTCTTAACCT
 AATAAATAAAAAAATACCAATTCTGAAAATTATAAAATACCTGATCTGGTAAATTGATGATTTGAAGATCTT
 AAAATCTGGAGCAAAGACTATTAAAGTAAGAAAATATTAAATCGAAGATTAACTCGACTAATAAAAGATGCAA
 AAAATTTGGGATTGAAATTAAATCAAATCTGCTTACAGAACGCAAGAATATCAAATTTTATTGATTACAA
 TGTCAAAACTTATGGCAGAAAAGTGCAGAAACCCAATCAGCAATTCCAGGCCATTCTCAACATCATGGAAC
 GCAATAGATTTATAAATATAGATGATAATTACTAAACACAAAAGAAGGAAATGGCTTATGAAAACCTCTAA
 AATACGGATTTCCGTTCATACCCAAAAGGATATGAAACGGACACTGGATATAAGCAGAGCCTGGCACTACTT
 ATACATAGGACCTAACCCATGCTTATTCAAGAAAATATTTAATAATTACAACATAAGCTTCTGAATTGG
 AACCCAGAACAAAACAAATCTTATTAAACCTAATTGAAAATATGCAAACCTAA

f342.aa

MLYLGDNKAMRTKIIIMTIIILAPISGFSNSKESARGKFGAGIILPLPIALQINIGNFDLDIGLYSGVNNLFSDW
 KTLFIALDYIFYIYTFPGAANILDFTSVGAGGYGTIWFSRGGSKSGSGPMSIGARLPLALNIAVFRKKFDIFLRIA
 PGLGMNVWSNGVGRWEVFAGLGLRFWFTZ

t342.aa

LAPISGFSNSKESARGKFGAGIILPLPIALQINIGNFDLDIGLYSGVNNLFSDWKTFLFIALDYIFYIYTFPGAANI
 LDFTSVGAGGYGTIWFSRGGSKSGSGPMSIGARLPLALNIAVFRKKFDIFLRIA PGLGMNVWSNGVGRWEVFAGL
 GLRFWFTZ

f342.nt

ATGCTATACTTAGGAGATAATAAGCAATGAGAACAAAATAATTATTATGACAATTATTATTAGCCCCAA
 TCTCAGGATTTCTAATTCAAAGAAATCTGCAAGGGTAAATTGGAGCAGGAATTATACTTCCATTACCAATTGC
 TCTACAGATTAATATAGGAAACTTGATCTTGACATTGGCTTACAGCGGAGTAAATAATTGTTTCAGACTGG
 AAAACATTATTATAGCATTAGACTATATTCTACATATAACACATTCCCGGGAGCTGCTAATATTGGATT
 CAGTTGGCGCAGGGGATATGGAACAATATGGTTCAAGATTGGAGGCAGTAAGTCAGGCTCAGGACCAATGAG
 CATTGGAGCAAGATTGCTTGGCTTAAATATTGAGTATTAGGAAGAAATTGACATATTTCAGAATAGCA
 CCCGACTTGGAAATGAATGTTGGAGTAATGGCGTTGGATTAGATGGAAAGTATTGAGGACTAAGAT
 TCTGGTTACTTAA

t342.nt

TTAGCCCCAATCTCAGGATTCTAATTCAAAAGAATCTGCAAGGGTAAATTGGAGCAGGAATTATACTTCCAT
 TACCAATTGCTCTACAGATTAATATAGGAAACTTGATCTTGACATTGGCTTACAGCGGAGTAAATAATTGTT
 TTCAGACTGGAAAACATTATTATAGCATTAGACTATATTCTACATATAACACATTCCGGAGCTGCTAATATT
 TTGGATTTCAGTTGGCGCAGGGGATATGGAACAATATGGTTCAAGATTGGAGGCAGTAAGTCAGGCTCAG
 GACCAATGAGCATTGGAGCAAGATTGCGTTGGCTTAAATATTGAGTATTAGGAAGAAATTGACATATT

TABLE 1. Nucleotide and Amino Acid Sequences

ACGAATAGCACCCGGACTTGGAAATGAATGTTGGAGTAATGGCGTTGGATTAGATGGAAAGTATTCGCAGGATTG
GGACTAAGATTCTGGTTACTTAA

f352.aa

MNKTKNRSLTYFIILSCISLFGANNNTISYSSIEIPLEDLSEEFKSSGNKSDQINTSKHLNKNIVSYEDPKKGKDL
KLPEINRDKKLPQKRMDEDNLKSVIENYENKNIKLLKTKNQKTSENEKKIESIEKKAKKYEILTNKLKNEIV
EIKKLLNKKIKPKEDENEKINIEEETDDDFEDNYEYNDEIEEQMRTITLLMKEZ

t352.aa

CISLFGANNNTISYSSIEIPLEDLSEEFKSSGNKSDQINTSKHLNKNIVSYEDPKKGKDLKLPEINRDKKLPQKRM
DENDLKSVIENYENKNIKLLKTKNQKTSENEKKIESIEKKAKKYEILTNKLKNEIVEIKKLLNKKIKPKED
NYEKINIEEETDDDFEDNYEYNDEIEEQMRTITLLMKEZ

f352.nt

ATGAATAAAACAAAAATCGAACGCCTTACGTATTTATAATACCTTCATGTATATCATTATTTGGGGCTAATAATA
ATACAATAAGCTACTCTAGCATTGAAATTCTCTAGAACACTTAAGTGAAGAACATTAAAAGTTCTGGAAATAAAAG
CGATCAAATAAAATACCTCAAAACATTAAACAAAAACATAGTTCTTATGAAGACCCAAAAAGGGTAAAGATCTA
AAATTGCCAGAAATATAAGAGACAAAAACTACCCAAAAAGAACATGGACAAAATGATCTAAATCTGTAATTG
AAAATTATGAAAATAAAATTAAAAACATAGAAAAGCTTTAAAACAAAAATCAAAAACATCGGAAATGAAAA
TAAAAAAATAGAATCAATCGAAAAAAAGCAAAAAATGAAATTAAACCAATAATTAAAAACGAAATAGTA
GAAATAAAAAGCTCTTAACAAAAAAATCAAGCCTAAAGAACATGAAAATTACGAAAAATAAATTGAAAACA
TTGAAGAAGAAACTGATGATGATTTGAAGACAATTATGAATATAATGATGAAATTGAAGAACAAATGAGGACAAT
TACCCCTCTAATGAAGGAATAA

t352.nt

TGTATATCATTATTTGGGGCTAATAATAACAAATAAGCTACTCTAGCATTGAAATTCTCTAGAACACTTAAGTG
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TGAAGACCCAAAAAGGGTAAAGATCTAAATTGCCAGAAAATATAAGAGACAAAAAAACTACCCAAAAAGAACATG
GACGAAAATGATCTAAATCTGTAATTGAAAATTATGAAAATAAAATTAAAACATAGAAAAGCTTTAAAACCA
AAAATCAAACATCGGAAATGAAAATAAAATAGAATCAATCGAAAAAAAGCAAAAAATATGAAATTTT
AACCAATAATTAAAAACGAAATAGTAGAAATAAAAAGCTCTTAACAAAAAAATCAAGCCTAAAGAACATGAA
AATTACGAAAAATAAATTGAAAACATTGAAGAACACTGATGATGATTTGAAGACAATTATGAATATAATG
ATGAAATTGAAGAACAAATGAGGACAATTACCCCTCTAATGAAGGAATAA

f301.aa

MQIDGKIYSIISFPVRDSVSTLGVIGILICFDESLDIIEENQYSSLKFGSKNYNFFMLDRNYMPIFSNLNNLQAKS
FSTAYSENFLSKVIAYAKKDSSSQYTFNERYDFYSLNFVKTDDFLTQGLILNVNSIPIMFKSNWVIFVAFLLSF
AIIIFYLCNTFVFSLINDFNRIVDYQKSKEKFKYSEDLN
EYLEQIETASNTESIDSSILVYEQLRDTFSRFEKSIVDILKGFEISIADPINDHNKYISEISSNFEESVSSFFYSID
KNLEIFNKVATINSTDIENIKSKVFDLNIVFENVNKNFADLLSQTNSLQSVNKLVLVSISAQTNMLAMNAAIEAAKA
GDAGKSFAVVAAEIRKLAINGKYSKTIKDELKTVDSIIAVINSEIDTIYKNFIDIQDNVDNNFSRHEKVDLTLAK
HFKEIGEFKERYLSDTKIRDAKNMYKEIFNNHYFISGKFNNFSQDLKEFKVSKMNLDAVSSLQEYSSLVKSSKDK
ILKTKEIQLQKINDEIKDILFZ

t301.aa

CFDESLDIIEENQYSSLKFGSKNYNFFMLDRNYMPIFSNLNNLQAKSFSTAYSENFLSKVIAYAKKDSSSQYTFN
YERDFYSLNFVKTDDFLTQGLILNVNSIPIMFKSNWVIFVAFLLSFAIIIFYLCNTFVFSLINDFNRIVDYQKS
DPFSLESPLEVYSSSIISYISSKLDNLSSKSNESFEKFKYSEDLN
EYLEQIETASNTESIDSSILVYEQLRDTFSRFEKSIVDILKGFEISIADPINDHNKYISEISSNFEESVSSFFYSID
KNLEIFNKVATINSTDIENIKSKVFDLNIVFENVNKNFADLLSQTNSLQSVNKLVLVSISAQTNMLAMNAAIEAAK
GDAGKSFAVVAAEIRKLAINGKYSKTIKDELKTVDSIIAVINSEIDTIYKNFIDIQDNVDNNFSRHEKVDLTLAK
HFKEIGEFKERYLSDTKIRDAKNMYKEIFNNHYFISGKFNNFSQDLKEFKVSKMNLDAVSSLQEYSSLVKSSKDK

TABLE 1. Nucleotide and Amino Acid Sequences

DELKTVDSIIAVINSEIDTIYKNFIDIQDNVDNNFSRHEKVDLTLAKHFKEIGEFKERYLSHDTKIRDAKNMYKEI
FNNHYFISGKFNNFSQDLKEFKVSKMNLDAVSSLQEYSSLVKSSKDKILKTKELIQKINDEIKDILFZ

f301.nt

ATGCAAATAGATGGAAAATTATTCTATAATAAGTTCCAGTTAGAGATTCTGTTCAACATTGGGTGTGATAG
GGATTTAATATGCTTGTAGAGTCGTTAGATATTGAAATCAGTTGATTCTCTAAATTGGTAGTAA
AAATTATAATTCTTGTGACAGAAATTACATGCCATTTCACAAACCTTAATAATCTCAGGCCAAATCT
TTTCTACAGCTTATAGTGAGAATTGGAGTAAAGTTAGCTTATGCTAAAAAGATTCTCTAGCTCTCAGT
ACACTTTAATTATGAAAGAGATTGGTAAACTTGTAAAAACCGATGATTGGACTCAGGGCTTAT
TTAAATGCAATTCCATTCTTAAATCAAATTGGTTATTTGTTGCATTAAATTGCTTAAATTGCTTT
GCAATTATTCTTATTCAGCAACTTTGTTCTTCAATTAAATGATTTAACAGAATTGTTGACTATCAA
AAATCAAAAGCGATCCTTGTAGCTTGAATCTCCCTAGAGGTTAAGTATTCTCATCTATTAAATTCTTAA
TCAGCTAGATAATCTGCTCTAAGAGTAATGATCTTGTGAGAAGATAAAATTAAATTCTGAAGATTGAAT
GAATATTGGAACAAATAGAAACTGCTATACAAATACTGAGAGTATAGATTCTAGCATTTAGTTACGAACAAAC
TAAGAGATACTTTCTAGATTGAAAATCAATTGTTGATATTAAAGGCTTGAATCTATTGCTGATCCGAT
TAATGATCACAATAAAATATCAGAAATCTTCAAAATTGGAGAGTGTAGTTCTTATAGTATAGAT
AAAAATTAGAAATTAAATTAGGTTGCTACTATAATTCTACTGATATTGAAATATTAAAGTAAGGTTTG
ATTAAATATTGTTGAAAATGTGAATAAAATTGAGCTTCAAGACATATGCTTGTATGAATGCAGCAATTGAAGCAGCAAAGCA
GGTGTGAGGTTAAAGTTGAGTTGCTGAGGAGATTAGAAAGCTGCTATTAAATTCTGAAATATTCTA
AAACCATTAAGATGAACCTAAACGGTCGACAGCATTATTGCACTATTAAATTGAGATTGATACAATTATAA
AAATTCTAGACATTCAAGATAATGTGGACAACAATTTCAGACACAGAGAAAGTAGATCTTACTCTGCTAAG
CATTAAAGAAATTGGCGAGTTAAAGAAAGGTATTGCTCACGATAACTAAGATCAGAGATGCTAAGAATATGT
ATAAAAGAAATATTAAATAATCATTATTATTAGTGGCAAGTTAACAACTTGTCAAGATTAAAGAGTTAA
AGTTCTAAGATGAATTAGTGCCTAAGTTCTCTCAAGAAATTCTCATTTAGTAAAGTCTTAAGGATAAG
ATATTAAAGACAAAGGAATTGATTCAAAAGATTAATGAGATTAAAGATATTCTTTAG

t301.nt

TGCTTGTAGTCGTTAGATATTGAAAATCAGTTGATTCTCTAAATTGGTAGTAAAATTATAATT
TTTTATGCTTGACAGAAATTACATGCCATTTCACAAACCTTAATAATCTCAGGCCAAATCTTCTACAGC
TTATAGTGAGAATTGGTGTAGTAAAGTTAGCTTATGCTAAAAAGATTCTCTAGCTCTCAGTACACTTTAAT
TATGAAAGAGATTCTTATTCTTAAACTTGTAAAACCGATGATTGGACTCAGGGCTTATTAAATGTCA
ATTCCATTCTTATTATGTTAAATCAAATTGGTTATATTGTTGCTATTAAATTGTTGCAATTATT
TTATTGCAATACTTTGTTCTTCAATTAAATGATTAAACAGAATTGTTGACTATCAAAATCAAAGC
GATCCTTTAGTCTGAATCTCCCTAGAGGTTAAGTATTCTCATCTATTATTCTTATATTGCTAACAGCTAG
ATAATCTGCTCTAAGAGTAATGAAATCTTGTGAGAAGATAAAATTCTGAAGATTGAATGAATATTGGA
ACAAATAGAAACTGCTATATCAAATACTGAGAGTATAGATTCTAGCATTTAGTTACGAACAACAAAGAGATACT
TTTCTAGATTGAAAATCAATTGTTGATATTAAAGGCTTGAATCTATTGCTGATCCGATTAATGATCACA
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AAATTAAATAAGGTTGCTACTATAAAATTCTACTGATATTGAAATATTAAAGTAAGGTTTGATTTAAATT
GTTTTGAAAATGTGAATAAAATTGAGCTTCAACAAATAGTTGCAAAAGTGTAAATAACTTT
TAGTTCAATTCTCAGACCAATATGCTTGTATGAATGCAGCAATTGAAGCAGCAAAGCAGGTGATGCAGG
TAAAAGTTGAGTTAAAGAAAGGTATTGCTCAGGAGATTAGAAAGCTTGTCTTAAATTCTGAAATATTCTAAACCAATTAA
GATGAACCTAAACGGTCGACAGCATTATTGAGCTTAAATTGAGATTGATACAATTATAAAATTCTAG
ACATTCAAGATAATGTTGACAACAATTCTCAAGACACAGAGAAAGTAGATCTTACTCTGCTAACGATTTAAAGA
AATTGGCGAGTTAAAGAAAGGTATTGCTCAGGAGATTAGAAAGCTTGTCTTAAATTGAGATTGCTAAGAATATGTATAAAGAAATA
TTAATAATCATTATTATTAGTGGCAAGTTAACAACTTGTCAAGATTAAAGAGTTAAAGTCTTAAGGATAAGATATTAAAGAC
AAAGGAATTGATTCAAAAGATTAATGAGATTAAAGATATTCTTTAG

f346.aa

TABLE 1. Nucleotide and Amino Acid Sequences

MSIDKVPDEAFAEKIVGDGIAILPTSNELLAPCDGKIGKIFKTNHAFSLETKEGVEIFVHFGINTLNLngKGFTRV
AEEGINVKQGEVIIRLDLEYLKEHSESVITPVVIANSDEVSSIEYSFGRENDSEYILSSSTVLTEEIRHKISQT
PVIAGKDLVLRVKKZ

t346.aa

CDGKIGKIFKTNHAFSLETKEGVEIFVHFGINTLNLngKGFTRVAAEGINVKQGEVIIRLDLEYLKEHSESVITPV
VIANSDEVSSIEYSFGRENDSEYILSSSTVLTEEIRHKISQT
PVIAGKDLVLRVKKZ

f346.nt

ATGTCAATTGATAAGGTTCCCGATGAAGCTTTGCTGAAAAAATAGTTGGCGATGGAATTGCAATTCTCCAACAA
GCAATGAGTTGGCGCCTTGTGATGGGAAAATAGGTAAAATTTTAAACCAATCATGCCTTAGCCTGAAAC
TAAAGAGGGCGTTGAAATTTCATTTGCAATTAACTCTTAATTAAATGTAAGGGTTTACAAGAGTT
GCTGAAGAGGGCATTAAATGTTAACAAAGGTGAAGTTATTAGGCTTGAATCTGAATATTAAAGAGCATTAG
AATCCGTTATTACTCCGGTTGTATTGCAAATTCTGATGAAGTTCAAGTATAGAATATTCTTTGGAAGGCTG
AAATGATTCTGAATATATTATCATCTCAACTGTCTTGACAGAAGAAATTAGGCATAAAATATCTCAAACAAAG
CCTGTTATAGCGGGCAAAGATTGGTGTGCGAGTTAAAAGTAA

t346.nt

TGTGATGGAAAATAGTAAAATTAAACCAATCATGCCTTAGCCTGAAACTAAAGAGGGCGTTGAAATT
TTGTCCATTGAAATTAAACTCTTAATTAAATGGTAAGGGTTTACAAGAGTTGCTGAAGAGGGCATTAAATGT
TAAACAAGGTGAAGTTATTAGGCTTGAATATTAAAGAGCATTAGAATCCGTTATTACTCCGGTT
GTTATTGCAAATTCTGATGAAGTTCAAGTATAGAATATTCTTTGGAAGGCTGAAAATGATTCTGAATATATT
TATCATCTCAACTGTCTTGACAGAAGAAATTAGGCATAAAATATCTCAAACAAAGCCTGTTAGCGGGCAAAGA
TTTGGTCTGCGAGTTAAAAGTAA

f373.aa

MNYQRIKNYCKFTSVFLFFLFSCVSNELKLDQSLVKGKLVNGLRYYIYKNQTPKNAVNMGIVFNVGSLNEEDNERG
IAHYLEHMAFNGTKDYPGNSIVDVLKKFGMQFGADINAATSFDFTYYRLDLSDGNNKDEIDESINILRNWASQISF
MKEEIDLERNIIIIEEKKLGETYPGRIYEKMDKFLTSGSLYEFRSPIGLEEQILSFQPEDFKKFYRKWYRPELASVI
VVGIDIPIEEKKIKKQFVSWKNPTDKIKEVKVSLDVELKDKFLLLEDLEVGEPSLMFFKKEIINFVTKDDLLNA
IKKSLLAALFENRFSELKTAGVKQFKNVSNKDFFSFKSDNNTIVAKSISLNFNPDHNEGIQDFFYELERIRKFGF
TQGELEKVRQSFYKSLELRKKNINKTNSWAIFQDLIEIAINGSNKFDMNEYCDLSFQYLEKIDLKTINNLVGREFD
VKNCAIFYSYHGRAHPVLTLEDIDNLQKIALKRELKPYENSIEGKFFKSLDDKDIIRENEFENEISSFVLENGV
EVYFKYNDQKKGVIDFSATSWGLINEDLKLIPVLSFAPGVVSGSGYGDYSALQIEKYLSDKAVSLRVGVGAQESY
ISGSSDKDLETLFQLIYFTFKEPKIDDVSLQNAINNIKALIKSNENSSDYHFKAISKFLNNNDPRFEDTKDSDL
QYFTKENILSFYKKRFTYANNFKFVLLETQIFRQZ

t373.aa

CVSNELKLDQSLVKGKLVNGLRYYIYKNQTPKNAVNMGIVFNVGSLNEEDNERGIAHYLEHMAFNGTKDYPGNSIV
DVLKKFGMQFGADINAATSFDFTYYRLDLSDGNNKDEIDESINILRNWASQISF
MKEEIDLERNIIIIEEKKLGETYPGRIYEKMDKFLTSGSLYEFRSPIGLEEQILSFQPEDFKKFYRKWYRPELASVI
NPTDKIKEVKVSLDVELKDKFLLLEDLEVGEPSLMFFKKEIINFVTKDDLLNAIKSLLAALFENRFSELKTAGV
KQFKNVSNKDFFSFKSDNNTIVAKSISLNFNPDHNEGIQDFFYELERIRKFGFTQGELEKVRQSFYKSLELRKKN
INKTNSWAIFQDLIEIAINGSNKFDMNEYCDLSFQYLEKIDLKTINNLVGREFDVKNCAIFYSYHGRAHPVLTLED
IDNLQKIALKRELKPYENSIEGKFFKSLDDKDIIRENEFENEISSFVLENGVEVYFKYNDQKKGVIDFSATSWG
GLINEDLKLIPVLSFAPGVVSGSGYGDYSALQIEKYLSDKAVSLRVGVGAQESYISGSSDKDLETLFQLIYFTF
EPKIDDVSLQNAINNIKALIKSNENSSDYHFKAISKFLNNNDPRFEDTKDSDLQYFTKENILSFYKKRFTYANNF
KFVLLETQIFRQZ

f373.nt

TABLE 1. Nucleotide and Amino Acid Sequences

ATGAATTATCAAAGAATTAAAGAATTATTGTAAATTACAAGCGTTTCTATTTTTGTTCCGTGTTCTA
 ATGAGTTAAAGTTAGATCAAAGTTGGTAAAGGAAAACCTGTCAATGGCTAAGGTATTATATTATAAAATCA
 AACCCCAAAGAATGCCGTTAATATGGGATTGTTTAATGTGGCTCACTTAATGAAGAAGATAATGAGAGGGGA
 ATAGCGCATTATCTTGAACATATGGCTTTAATGGTACAAAAGATTATCCAGGGAAATTCTATAGTTGATGTTCTTA
 AAAAATTGGAAATGCAATTGGTGCTGACATTAATGCTGCTACTAGTTGATTCACTTATTATAGACTGATT
 GTCAGATGTAATAAAAGATGAAATTGATGAATCTATAAAATTGGAGAAACTGGGCTCTCAAATCAGTTTC
 ATGAAAGAAGAAATAGATCTAGAGCGAAATATTATTGAGGAAAAAAAGCTTGGTGGAGACTTATCCTGGAAGAA
 TTTATGAGAAAATGGATAAGTTTGACAAGCGGAAGTCTTATGAATTAGAAGTCTTATGGACTTGAAGAGCA
 AATTATCTTCAAGCCAGAAGATTAAAGGGATAGAAATTGAGAAGATAAAAGCAATTGGGAACTTGAAGTGTATT
 GTGGTAGGAGATATTGATCCTATAGAAATTGAAGAGATAAAAGCAATTGGTCTTGTGAAAGGAAATCCAACCG
 ATAAAATTAAAGAAGTAAAGTAAGTTAGCGTAGAGCTTAAGGATAAATTAACTTTAGAAGATTGGAAAGT
 TGGAGAGCCTAGTTAATGTTCTTAAAAGGAAATTATTAACCTTGTAAAGACCAAGATGACCTTAAATGCT
 ATTAAAAAGTCTTATTAGCCGCTCTTTGAAAATAGATTCTGAATTAAAGACTGCTGGGTAAGCAATTAA
 AAAATGTTCAAATAAAGATTCTCATTAAATCAGATAACAATACATTGTTGCAAAATGATTCTTAA
 CTTAATCCAGATCATTGAAACGAAGGAATACAAGACTTTTATGAGCTTGAGAGGATAAGAAAATTGGATT
 ACCCAAGGTGAGCTGAAAAAGTTAGATCTCAATTAACTTAAAGGAAATTGCTATTAAATGGTCTAATAAATTTGAT
 CAAATTCACTGGCTATTTCAGGATTTAATAGAAATTGCTATTAAATGGTCTAATAAATTTGAT
 TTGCGATCTTCTTCAATATTGGAAAAGATTGATTAAAACAATAACAATTCTGTAGGAAGAGAGTTGAT
 GTAAAAAATTGTGCAATTTCATTCTACATGGAAAGAGCACATCCTGTTAACTCTTGAAGATATTGACAATC
 TTCAAAAGATAGCTTAAAAGAGAGTTAAAGCCTTATGAGAATTCTTAAATTGAAGGTAATTGGGTTAAGAAGTC
 TTTAGATGATAAAAGATATTAGAGAAAATGAGTTGAAAATTGAAATTCTGTCAATTGTTCTGAAAATTGGGTT
 GAAGTTATTAAATATAATGATCAAAAAAGGTGTAATTGATTAGTCAACTTCTGGGAGGTTAATT
 ATGAAAGATTAAACCTATTCCGTTTATCTTGTCCCGAGTAGTATCTGGTTCGGGTTATGGTGATTATTC
 TGCATTACAGATTGAAAATATTATCAGATAAAAGCTTCTTAAAGAGTTGGGTTGGAGCTCAAGAATCATAT
 ATTCTGGAAGATTGAGATAAAAAGATCTGAAACTCTTTCACTTATATTTTACTTTAAGGAAACCCAAA
 TTGATGATGTTCTTGCAAAATGCTATTAAATAATAAAACATTAAATAAGAGCAATGAAAATAGTTGATTA
 TCATTTCATAAAAGCATTAGTAAATTAAACATAATGATCCTAGATTGAGATAACAAAGATAGTGAATT
 CAATATTTCATAAAAGAAAATATTGTCTTTATAAGAAAAGGTTACTATGCAAATAATTAAAGTTGAT
 TGCTGGAGACTCAGATATTCAAGACAATAA

t373 nt

TGTGTTCTAATGAGTTAAAGTTAGATCAAAGTTGGTAAAGGAAAACCTGTCAATGGCTAAGGTATTATTT
 ATAAAAATCAAACCCCAAAGAATGCCGTTAATATGGGATTGTTTAATGTGGCTCACTTAATGAAGAAGATAAA
 TGAGAGGGAAATAGCGCATTATCTTGAACATATGGCTTTAATGGTACAAAAGATTATCCAGGGAAATTCTATAGTT
 GATGTTCTTAAAAATTGGAAATGCAATTGGTGCTGACATTAATGCTGCTACTAGTTGATTCACTTATTATA
 GACTTGATTTGTCACTGGTAATAATAAAGATGAAATTGATGAATCTATAAAATTGAGGAAACTGGCTCTCA
 ATCAGTTCTATGAAAGAAGAAATAGATCTAGAGCGAAATATTATTGAGGAAAAAAAGCTTGGTGGAGACTTAT
 CCTGGAAGAATTATGAGAAAATGGATAAGTTTGACAAGCGGAAGTCTTATGAATTAGAAGTCTTATTGGAC
 TTGAAGAGCAAATTATCTTCACTGGCAGAAGATTAAAGGAAATTGAGGAAATTGAGGAAATTGAGGAA
 AAGTGTATTGTGGTAGGAGATTGATCCTATAGAAATTGAGGAGATAAAAGAAGCAATTGTTCTGGAAA
 AATCCAACCGATAAAATTAAAGAAGTAAAGTAAGTTAGACGTTAGAGCTTAAGGATAAATTGAGGAA
 ATTGGAAGTTGGAGAGCCTAGTTAATGTTCTTAAAGGAAATTATAACTTGTAAAGACCAAAGATGACCT
 TTTAAATGCTATTAAAGTCTTATTAGCCGCTCTTTGAAAATAGATTCTGAAATTAAAGACTGCTGGGTA
 AAGCAATTAAAGATTGTTCAAATAAAAGATTCTCATTTAAATCAGATAACAATACATTGTTGCAAAATCGA
 TTTCTTAACTTAAATCCAGATCATTGAAACGAAGGAATACAAGACTTTTTATGAGCTTGAGAGGATAAGAAA
 ATTGGAATTACCAAGGTGAGCTTGAAGGAAAGTTAGATCTCAATTAAACAAATCTTGAATTAAAGGAAAAGAAT
 ATAAATAAAACAAATTGCGATCTTCTTCAATATTGGAAAAGATTGATTAAAACAATAACAAATCTGTAGGAAG
 AGAGTTGATGTAAGGAAATTGTGCAATTCTTATTCTTACATGGAAAGAGCAGCACATCCTGTTAACTCTTGAAGAT
 ATTGACAATCTCATAAAAGATAGCTTAAAGAGAGTTAAAGCCTTATGAGAATTCTTAAATTGAAGGTAATT
 TTAAGAAGTCTTATTAGATGATAAAAGATATTAGAGAAAATGAGTTGAAAATTGAAATTTCGTCAATTGTTCTG
 AAATGGGTTGAAGTTATTAAATATAATGATCAAAAAAAAGGTGTAATTGATTAGTCAACTTCTGGGGA
 GTTTAATTAAATGAAGATTAAACCTATTCCGTATTCTTATCTTGTCCCGAGTAGTATCTGGTTCGGGTTATG
 GTGATTATTCTGCATACAGATTGAAAAAATTATCAGATAAAAGCTGTTCTTAAAGAGTTGGGTTGGAGCTCA
 AGAATCATATATTCTGGAAGTTGAGATAAAAGATCTGAAACTCTTTCACTTGTATATATTACTTTAAG

TABLE 1. Nucleotide and Amino Acid Sequences

GAACCCAAAATTGATGATGTTCTTGCAAAATGCTATTAAATAATATAAAAGCATTAAATAAGAGCAATGAAAATA
GTTCTGATTATCATTTCATAAAGCCATTAGTAAATTTAAACAATAATGATCCTAGATTTGAAGATAACAAAAGA
TAGTGATTGCAATATTTACAAAAGAAAATTTGTCTTTATAAGAAAAGCTTACTTATGCAAATAATTT
AAAGTTGCTTGCTGGAGACTCAGATATTCAAGACAATAA

f384.aa

MDWDFEKIIFLNESTRLALSGCAKLILDFKSDGSIVTQVDKQIEQFLFKEIKKPGNFVLGEETISTYKEEYIKDA
LISESTFIIDPIDGTSSFAAGLPSYGISLAYASGGKIIIEGAISLPLSGEFFITSKDNVFYAKKNIGSYPLKDFNK
FIFDNSKCYNIHSLLAVRSIIIRLFNLDISSHIHINGSCVYSFAKLFTGSYKAYFSFVGLWDIAACLAIGNKLGMV
GEFYCGNKMTLDILDMSYILEPNNHKRWSLKDFFIYSDNKSTIDIIRKDANKKINK

t384.aa

CAKLILDFKSDGSIVTQVDKQIEQFLFKEIKKPGNFVLGEETISTYKEEYIKDALISESTFIIDPIDGTSSFAAGL
PSYGISLAYASGGKIIIEGAISLPLSGEFFITSKDNVFYAKKNIGSYPLKDFNKFIFDNSKCYNIHSLLAVRSII
RLFNLDISSHIHINGSCVYSFAKLFTGSYKAYFSFVGLWDIAACLAIGNKLGMVGEFYCGNKMTLDILDMSYILEP
NNHKRWSLKDFFIYSDNKSTIDIIRKDANKKINKZ

f384.nt

ATGGATTGGGATTTGAAAAAAATATATTTATTAAATGAATCAACTAGGCTTGCAATTAGGGTTGTGCTAAAT
TAATTTAGATTTAAATCTGATGGGTCTATTGTAACTCAGGGTGTAAAGCAAATTGAGCAATTCTTATTCAAAGA
GATCAAAAAGCCTGGAAATTTGTTCTGGAGAAGAGACAATATCTACTTATAAAGAAGAGTATATCAAAGATGCT
TTAATATCAGAGAGTACTTTATTGATCCTATTGATGGAACTTCTTGTGCTTTGAGCAGGCCCTCCTCATATG
GAATATCGCTAGCGTATGCTAGTGGCGGAAAATTATTGAGGAGCCATTCTCTCCCTTAAGCGGAGAGTTTT
TATTACTCTAAAGATAATGTATTTATGCTAAAAAAACATTGGTAGCTATCCTTAAAGGATTTAATAAA
TTTATTTGATAATTCTAAATGTTACAATATTGATAGTTACTGCAAGTCTATTATAAGGTATTTA
ATCTGATATTCTCTCATATTGATATTGATATTGTTCTGTGTATATTCTTGTAAACTTTTACAGGTTCTTA
TAAGGCCTACTTTCTTGAGACTTGGGATATTGCAAGCTGTTAGCTATTGTAATAAAATTGGGATGGTT
GGCAATTTTATTGTTGAGACTTGGGATATTGCAAGCTGTTAGCTATTGTAATAAAATTGGGATGGTT
AAAGATGGCCTTGAAAGATTTTATTCTGATAATAATCAACAATAGACATTATAAGAAAAGATGCAAA
AAAAAAATCAATAAGTAA

t384.nt

AGTGGTTGTCTAAATTAGATTTAAATCTGATGGGTCTATTGTAACTCAGGTTGATAAGCAAATTGAGC
AATTCTTATTCAAAGAGATCAAAAGCCTGGAAATTTGTTCTGGAGAAGAGACAATATCTACTTATAAAGAAGA
GTATATCAAAGATGCTTAAATATCAGAGAGTACTTTATTGATCCTATTGATGGAACTTCTTCTTGCAGCA
GGCCTCCTCATATGGAATATCGCTAGCGTATGCTAGTGGCGGAAAATTATTGAGGAGCCATTCTCTTCTT
TAAGCGGAGAGTTTTATTACTCTAAAGATAATGTATTGCTAAACATTGGTAGCTATCCTTAA
AAAGGATTTAATAAATTATTGATAATTCTAAATGTTACAATATTGATAGTTACTGCAAGTCTCAAGGTCT
ATTATAAGGTTATTAAATCTGATATTCTCTCATATTGATATTGTTCTGTGTATATTCTTGTAAAC
TTTTACAGGTTCTTATAAGGCTACTTTCTTGTAGGACTTGGGATATTGCAAGCTGTTAGCTATTGGTAA
TAAATTGGGATGGTGGCAATTATTGTTGAGACTTGGGATATTGCAAGCTGTTAGCTATTGGTAA
GAGCCTAATAATCATAAAAGATGGCCTTGAAAGATTTTATTCTGATAATAATCAACAATAGACATT
TAAGAAAAGATGCAAATAAAATCAATAAGTAA

f860.aa

MAFYKLNDNIALAEDLLKYLLSSILNECSQDMDFLENYIEKGLIKKLENVINSNFevityTKAIEILENSKKNFEI
KPYWGIDLQTDHERYLTEETFKPVVVIDYPKNFKAFYMKANKDNKTVKGMDILVPKIGEIIGGSEREDDLQKLEN
RIKELNLNIEHLNWYLDLRRFGSAPHSGFGLGLERLVQYSTGISNIRDSDIPFPRTPKNLYFZ

t860.aa

TABLE 1. Nucleotide and Amino Acid Sequences

CSQDMDFLENYIEKGLIKKLENNVINSNFEVITYTKAIEILENSKKNFEIKPYWGIDLQTDHERYLTEETFKKPVVV
IDYPKNFKAFYMKANKDNKTVKGMDILVPKIGEIIIGGSEREDDLQKLENRIKELNLNIEHLDLRRFGSAPHS
GEGLGLERLVOYSTGISNIRDSSIPFPRTPKNLYFZ

- f860, nt

ATGGCTTTATAAGCTAACGACAATATTGCCCTAGCAGAAGATCTTGAAATATCTTTAAGTTCAATTAA
ACGAATGCTCACAAAGATATGGATTTAGAAAATTACATTGAAAAAGGTTAATTAAAAACTAGAAAATGTAAT
AAATTCAAATTTGAGGTTATTACCTATACTAAAGCAATTGAAATTCTGAAAACCTCAAAAAAAATTGGAAATA
AAACCTTACTGGGAATAGATTGCAAACAGATCAGAAAGATACTAACAGAAAGAGACTTTAAAAACCGGTAG
TGGTCATTGATTATCCAAAAATTCAAAGCATTTACATGAAAGCAAATAAGACAATAAAACTGTTAAAGGAAT
GGACATACTTGGTCCAAAAATTGGAGAGATTATAGGGGAAGCGAAAGAGAAGATGACCTCAAAAATTAGAAAAT
AGAATAAAAGAATTAAACTTAAACATTGAACATCTAAACTGGTATCTGATCTAACAGGAATATCTAATATAAGAGATTCAATACC
ATTCTGGCTTGAGCTTGGACTTGAAGAGATTGGTCAACTAACAGGAATATCTAATATAAGAGATTCAATACC
ATTCCCAAGGACTCCTAAATCTTATTAA

t860.nt

TGCTCACAAAGATATGGATTTAGAAAATTACATTGAAAAAGGTTAATTAAAAAACTAGAAAATGTAATAAATT
CAAATTGAGGTTATTACCTATACTAAAGCAATTGAAATTCTTGAAACTCAAAAAAAATTGAAATAAAACC
TTACTGGGAATAGATTGAAACAGATCACGAAAGATACCTAACAGAAGAGACTTTAAAAACCGGTAGTGGTC
ATTGATTATCCTAAAGCATTTCATGAAAGCAATAAGACAATAAAACTGTTAAAGGAATGGACAA
TACTTGTCCAAAATTGGAGAGATTAGGGGAAGCAGAAGAGAAGATGACCTCAAAATTAGAAAATAGAAT
AAAAGAATTAAACTAAACATTGAACATCTAAACTGGTATCTGATCTAAGAAGATTGGCTGGCTCCTCATTCT
GGCTTGGACTGGACTTGAAGATTGGTGCAATACTCACAGGAATATCTAATATAAGAGATTCAATACCATTC
CAAGGACTCCTAAATCTTATTGTTAA

f446.aa

MKILRLCFLFLFFACTFDYDEYSSRSVAKKFPSIQILGIKYYDVVYNKEQTVLNSLSFSYFNDYKIVKAENGRL
YHSLDNEISGKFNNLEGSYITKDLDMRDSVEFKIEDKNYYLLNSNRLLWKNDKKLQSPPNELVLIRFNDSKING
KGFSYFLKSNVFYFDSGVEGIMNZ

t446.aa

CTFDYDEYSSRSDVAKKFPSIQILGIKYYDVVYNKEQTVLNSLSFSYFNDYKIKYKAENGRLYHSLDNEISGKFNN
LEGSYITKDLDMRDSVEFKIEDKNNNYLLNSNRLLWKNKDQLQSPPNELVLRFDNSKINGKGFSYFLKSNVYF
DSGVEGIMNZ

f446.nt

ATGAAAATACCTAGCTTGTGTTGTTGTTGCTTGACTTTGATTATGATGAGTATTCTAGTAGAT
CTGATGTGGCCAAAAGTTCTTCAATACAAATATTAGAATCAAGTATTATGATGTTGATACAATAAAGAGCA
AACCGTTAAATTCTTAAGCTTAGTTAGTTCAATGACTATAAAATTATAAGGCAGAGAATGGAAGGTTTTA
TATCATTCCCTAGATAATGAAATTTCAGGGAAAGTTAATAATTGGAAGGTTCTTATATAACAAAGGATTGGATA
TGAGAGATTCTGTAGAATTAAAATAGAAGATAAAAATAATTATTATTTGCTTAATTCAAATAGGCTTTATGGAA
GAATAAAGACAAGAAGTGTCAATCCCCCCCCAAATGAGCTAGTTAATTAGATTAAATGATAGCAAATAAACCGGA
AAACGGATTTCCTATTAAAGAGCAATGTTTTATTGATTCTGGAGTTGAAGGAATCATGAATTGA

t446.nt

TGTACTTTGATTATGATGAGTATTCTAGTAGATCTGATGTCGCCAAAAAGTTCCCTCAATACAATATTAGGAA
TCAAGTATTATGATGTTGATACAATAAAGAGCAACCGTTTAAATTCTTAAGCTTAGTTATTCAATGACTA
TAAAATTATAAGGCAGAGAATGGAAGGTTTATATCATTCCCTAGATAATGAAATTTCAGGGAAAGTTAATAAT
TTGGGAAGGTTCTATATTACAAGGATTGGATATGAGAGATTCTGAGAATTAAAATAGAAGATAAAAATAATT
ATTATTTGCTTAATTCAAAATAGGCTTTATGGAAGAATAAGAGACAAGAGTTGCAATCCCCCCCCTAAATGAGCTAGT

TABLE 1. Nucleotide and Amino Acid Sequences

ATTAATTAGATTTAATGATAGCAAAATAACGGAAAAGGATTTCTTATTAAAGAGCAATGTTTTATT
GATTCTGGAGTGAAGGAATCATGAATTGA

f457.aa

MKQKLSWILLFCFLSCRSESRLAENVLIEFFDSIKNFQSSPEIFFNYLNIPSDDLKAKIRGLKSQAKDDFIFYPL
FFNNLRYEIIGRKNISKGFEEVVIKNINFQNGIEKFLAKLNKIEGRSLNIKLNLEKKERKKIFDNLINEVIGELDD
FDYTEVVHFFRKKSSSESYKIELLGDVLNQSRNKLINDLFLVSPGIZ

t457.aa

CFLSCRSESRLAENVLIEFFDSIKNFQSSPEIFFNYLNIPSDDLKAKIRGLKSQAKDDFIFYPLFFNNLRYEIIG
RKNISKGFEEVVIKNINFQNGIEKFLAKLNKIEGRSLNIKLNLEKKERKKIFDNLINEVIGELDDFDYTEVVHFFR
VVKSSSESYKIELLGDVLNQSRNKLINDLFLVSPGIZ

f457.nt

ATGAAGCAAAATTAAGTTGGATTATTATTTGTTCTGTAGATCTGAATCTAGATTGGCTGAAAATG
TTTAATAGAGTTTTGATTCTATTAAAATTTCAAAGCAGTCCTGAAATATTAAATTATTAATTTAAATTATTCC
AAAGTGTGATCTGAAGGCAAAATTCTGGGGTTGAAATCTCAGGCAAAGGATGATTCTAGTTTATCCTTTG
TTTTTAATAATCTAAGATATGAGATAATAGGTAGAAAAAATATTCTAAGGGCTTGAATTGAAAGTTGTTATT
AAAATTTAACTTCAAAACGGTATAGAAAAAATTTTGGCTAAATTAAATAAAATTGAAGGGAGATCTTAAATAT
AAAAAATTAGAAAAAAAGAGCGTAAAAAAATATTGACAATTAAATAATGAAGTTATTGGAGAGTTGATGAT
TTGATTACACTGAAGTTGTTCATTTTTAGACTAGTTAAGAGTTCTCTGAAAGTTATAAAATAGAGCTTTAG
GAGATGTTAAATATACAGTCTAGAAATAAGCTTATTATGATCTTTGGTTTATCGCTGGAATTAA

t457.nt

TGTTTTTGCTTGTAGATCTGAATCTAGATTGGCTGAAAATGTTAATAGAGTTTGATCTATTAAAATT
TTCAAAGCAGTCCTGAAATATTAAATTATTCAGTCAGGCAAAGGATGATTCTCATTTTATCCTTTGTTTAATAATCTAAGATATGAGATAATAGGT
GTTGAAATCTCAGGCAAAGGATGATTCTCATTTTATCCTTTGTTTAATAATCTAAGATATGAGATAATAGGT
AGAAAAAAATTTCTAAGGGCTTGAATTGAAAGTTGTTATTAAAATATTAACTTCAAAACGGTATAGAAAAAAT
TTTGGCTAAATTAAATAAAATTGAAGGGAGATCTTAAATATTAAAATTAGAAAAAAAGAGCGTAAAAAAAT
ATTGACAATTAAATAAAATGAAGTTATTGGAGAGTTGGATGATTCTGAAAGTTGTCATTAGA
CTAGTTAAGAGTTCTCTGAAAGTTATAAAATAGAGCTTTAGGAGATGTTAAATATACAGTCTAGAAATAAGC
TTATTAAATGATCTTTGGTTTATCGCTGGAATTAA

f542.aa

MRIVIFIFGILLTCSRNGIESSSKKIKISMLVDGVLDKSFNSSANEALLRLKKDFPENIEEVFSCAISGVYSS
YVSDLDNLKRNGSDLIWLVGYMLTDASLLVSSENPKISYGIIDPIYGDDVQIPENLIAVVFRVEPRCFFGWLYCSQ
KKLFWQNRFYRGNEGZ

t542.aa

CFSRNGIESSSKKIKISMLVDGVLDKSFNSSANEALLRLKKDFPENIEEVFSCAISGVYSSYVSDLDNLKRNGSD
LIWLVGYMLTDASLLVSSENPKISYGIIDPIYGDDVQIPENLIAVVFRVEPRCFFGWLYCSQKKLFWQNRFYRGNE
GZ

f542.nt

ATGAGAATTGTAATTAACTCGGTATTGACTTCTGCTTAGTAGAAATGGAATAGAAATCTAGTTCAA
AAAAAATTAAAGATATCCATGTTGGTAGATGGTGTCTGACGACAATCTTTAATTCTAGTCTAATGAGGCTTT
ATTACGCTGAAAAAAGATTTCCAGAAAATATTGAAGAAGTTTTCTGTGCTATTCTGGAGTTATTCTAGT
TATGTTCAAGATCTTGATAATTAAAAGGAATGGCTCAGACTTGATTTGGCTGTAGGGTACATGCTTACGGACG
CATCTTATTGGTTCATCGGAGAATCCAAAATTAGCTATGGAATAATAGATCCCATTATGGTGTGATGTTCA

TABLE 1. Nucleotide and Amino Acid Sequences

GATTCCCTGAAAACCTTGATTGCTGTTTCAGAGTAGAGCCAAGGTGCTTTTGCTGGCTATATTGCAGCCAA
AAAAAGCTTTCTGGCAAAATAGGTTTATAGGGGAATGAAGGGTAA

t542.nt

TGCTTTAGTAGAAATGGAATAGAATCTAGTCAAAAAAAATTAAGATATCCATGTTGGTAGATGGTGTCTGACG
ACAAATCTTTAATTCTAGTCTAATGAGGCTTATTACGCTGAAAAAAGATTTCAGAAATATTGAAGAAGT
TTTTCTTGCTATTCTGGAGTTATTCTAGTTATGTTACAGATCTGATAATTAAAAGGAATGGCTCAGAC
TTGATTGGCTTGAGGGTACATGCTACGGACGCATCTTATTGGTTCATCGGAGAATCCAAAATAGCTATG
GAATAATAGATCCCATTATGGTGATGATGTTACAGATTCTGAAACATTGATTGCTGTTTCAGAGTAGAGCC
AAGGTGCTTTTGCTGGCTATATTGCAGCCAAAAAAGCTTTCTGGCAAAATAGGTTTATAGGGGAATGAA
GGGTAA

f93.aa

MKRILAMHDISSMGRSLTICIPVISSFMQVCVPFTAVLSASTAYKKFEIVDLTDHLEKFINIWEQNEHFDILY
TGFLGSEKQQITIEKIIKLIKFEKIVIDPVFADDGEIYPIFDNKIIISGFRKIIKYANIITPNITELEMLSKSSKLN
NKDDIIKAILNLDTKATVVVTSVKGRLGNLGNICYNPKNKEYSEFFLEGLEQNFSGTGDLFTSLIGYLEKFETEQA
LEKTTKAIHLIILKESIKENVSKKEGVRIENFLKNTFZ

t93.aa

CIPVISSFMQVCVPFTAVLSASTAYKKFEIVDLTDHLEKFINIWEQNEHFDILYTGFLGSEKQQITIEKIIKLI
KFEKIVIDPVFADDGEIYPIFDNKIIISGFRKIIKYANIITPNITELEMLSKSSKLNKDDIIKAILNLDTKATVVV
TSVKGRLGNLGNICYNPKNKEYSEFFLEGLEQNFSGTGDLFTSLIGYLEKFETEQA LEKTTKAIHLIILKESIKENV
SKKEGVRIENFLKNTFZ

f93.nt

ATGAAAAGAATTTAGCAATGCATGATATTCAAGCATGGGAAGAACATCTCTTACAATATGCATACCAAGTAATAT
CTTCGTTTAATATGCAAGTTGCTCTTGTGACAGCTGCTTCTGCTTCCACAGCTTATAAAAATTGAAAT
AGTGGATTAAACGATCATTAGAAAATTATCAATATATGAAAGAACAAAATGAGCACTTGTACATACTCTAT
ACCGGATTCTGGGAAGCGAAAACAACAAATAACAATAGAGAAAATAATTAAATAAAAATTGAAAATTG
TAATTGATCCTGTGTTGCTGACGATGGAGAAATTACCTATATTGATAATAAAATAATTAGTGGATTAGAAA
AATCATAAAAGTACGCAAACATAATAACACCCAATATCACAGAACTTGAATGCTAAGCAAAGCTAAAACTTAA
ACAAAGATGATATCATAAAAGCAATATTAAATCTGATACAAAAGCGACGGTAGTTGTTACAAGCGTAAAAGGG
GAAATCTCTGGAAACATTGCTACAATCCTAAAACAAGAATACTCGGAGTTTTTTAGAAGGATTAGAACA
AAATTTCAGTGGAACAGGAGATTATTACAGCTACTTATAGGATATTGGAAAATTGAAACAGACCAAGCC
TTAGAAAAACAAACAAAGGCTATTACCTAATAATAAAAGAGTCATTAAAGAAAATGTTCAAAAAAGAAGGG
TCCGAATTGAAAATTCTTAAAAATACATTGAA

t93.nt

TGCATACCACTAATCTCGTTAATATGCAAGTTGCTCTTGTGACAGCTGCTTCTGCTTCCACAGCTT
ATAAAAAAATTGAAATAGTGGATTAAACCGATCATTTAGAAAATTATCAATATATGAAAGAACAAAATGAGCA
CTTGACATACTCTATACCGGATTCTGGGAAGCGAAAACAACAAATAACAATAGAGAAAATAATTAAATA
AAATTGAAAAATTGTAATTGATCCTGTGTTGCTGACGATGGAGAAATTACCTATATTGATAATAAAAATA
TTAGTGGATTAGAAAATCATAAGTACGCAAACATAAAACACCCAATATCACAGAACTTGAATGCTAAGCAA
AAGCTAAAACCTAACAAACAAAGATGATATCATAAAAGCAATATTAAATCTGATACAAAAGCGACGGTAGTTGTT
ACAAGCGTAAAAGGGAAATCTCTGGAAACATTGCTACAATCCTAAAACAAGAATACTCGGAGTTTTTT
TAGAAGGATTAGAACAAAATTCACTGGAACAGGAGATTATTACAGCTACTTATAGGATATTGGAAAATT
TGAAACAGAGCAAGCCTAGAAAAACAAACAAAGGTATTACACCTAATAATAAAAGAGTCATTAAAGAAAATGTT
TCAAAAAAGAAGGGTCCGAATTGAAAATTCTTAAAAATACATTGAA

f105.aa

TABLE 1. Nucleotide and Amino Acid Sequences

MGLYLKLLRQSINLKS LPLSVLFFSCNVVDTDFS VLEFKVANFNLNDDFSQGLLDSAYNILNRSFDLIIIKNLKN
 KNVLDLINNRVLFR AFKNAYFIDQGSGLSVSILSKRKINIKVLSVMQDSCDLKLGLLVDFKFENNHYGIVIYNLSK
 DFIKSIANLQISEQILYLKAQMDKLMFILDESEFVIFDLLIKNGFFSLINDSNYTSMLANKIDFRVFSNFFARVSL
 YSFMFVIADYLHSNYVVENFPQKIVINZ

t105.aa

CNVVDTDFS VLEFKVANFNLNDDFSQGLLDSAYNILNRSFDLIIIKNLKNKNVLDLINNRVLFR AFKNAYFIDQGS
 GLSVSILSKRKINIKVLSVMQDSCDLKLGLLVDFKFENNHYGIVIYNLSKDFIKSIANLQISEQILYLKAQMDKLM
 FILDESEFVIFDLLIKNGFFSLINDSNYTSMLANKIDFRVFSNFFARVSLYSFMFVIADYLHSNYVVENFPQKIVI
 NZ

f105.nt

ATGGGCTGTATTGAGACAAAGTATCAACTTGAAGAGTTATTCCGCTTAGTGT TATT TTT
 CCTGTAATGTTGAGATA CAGATTAGTGTGTTAGGTTAAGGTTGCAAA TTTAATTTAAATGATGATTTC
 TCAAGGGTTACTTGATTCTGCTTATAATATTCTAAATCGAAGTTGATTAAATAATTAAAGAATCTTAAAGAAT
 AAAAATGTTCTGATTAAATAATAGAGTTTATTAAGACCTTTAAGAATGCTTATTGATCAAGGTA
 GTGGCCTTCTGTTAGCATTCTCTAAAGCGCAAAATAATTAAGGTTAAGTGTAAAGATTCTTCAAGGTA
 TTTAAATAGGATTGCTTGTGGATTAAATTGAGAATAATCACTATGGTATTGTTATTATAATTAAAGCAAG
 GATTTATTAAAGTATTGCCAATTGCAAATTAGTGAACAAATTATTAAGGCCAAATGGATAAATTGA
 TGTTTATTAGATGAATCTGAATTGTTATTGATTAAATCAAATGGATTAGCTTAAATAATGA
 TTCAAACTACACTCAATGTTAGCAAATAAAATTGATTAGTTCTAATTGCTAGGGTTCTTTA
 TATTCAATTGTTGTAATTGCAGATTATTGCATAGCAATTATGTTGAGAATTTCCTCAAAATAGTTA
 TCAATTGA

t105.nt

TGTAATGTTGAGATA CAGATTAGTGTGTTAGGTTAAGGTTGCAAA TTTAATTAAATGATGATTTC
 AAGGGTTACTTGATTCTGCTTATAATATTCTAAATCGAAGTTGATTAAATAATTAAAGAATCTTAAAGAATAA
 AAATGTTCTGATTAAATAATAGAGTTTATTAGAGCTTTAAGAATGCTTATTGATCAAGGTAGT
 GCCCTTCTGTTAGCATTCTCTAAAGCGCAAAATAATTAAGGTTAAGTGTAAAGTCAAGATTCTTGC
 TAAATAGGATTGCTTGTGGATTAAATTGAGAATAATCACTATGGTATTGTTATTAAAGCAAGGA
 TTTTATTAAAGTATTGCCAATTGCAAATTAGTGAACAAATTATTAAGGTTAAGCTTAAATAATGATT
 TTTATTAGATGAATCTGAATTGTTATTGATTAAATCAAATGGATTAGCTTAAATAATGATT
 CAAACTACACTCAATGTTAGCAAATAAAATTGATTAGTTCTAATTGCTAGGGTTCTTTA
 TTCATTATGTTGTAATTGCAGATTATTGCATAGCAATTATGTTGAGAATTTCCTCAAAATAGTTA
 AATTGA

f150.aa

MKTFVIIGLSNLGIHLLLEDLSRLDCQIIIDTSKELIEEYDVISTESFVVEQFTKNALKRIIPVDTDAVVIDFDDD
 LGKSALVTHYCNLLGLKEICVKTENRDDAEILKTLGATKIIIFPSKDAARRLTPLLSPNLSTYNIIGYDIIVAEV
 IPKEYVGKTLFEADLRRECGITVIAVRNLSNSRYEFVGDYFFLKDDKIVICGKPDSIENFTNNKDLIKDLISGSK
 EDENLNKDAEKKSRFLGIFNFMKIFQKDRKDNZ

t150.aa

CQIIIDTSKELIEEYDVISTESFVVEQFTKNALKRIIPVDTDAVVIDFDDDLGKSALVTHYCNLLGLKEICVKTE
 NRDDAEILKTLGATKIIIFPSKDAARRLTPLLSPNLSTYNIIGYDIIVAEVTPKEYVGKTLFEADLRRECGITVI
 AVRNLSNSRYEFVGDYFFLKDDKIVICGKPDSIENFTNNKDLIKDLISGSKDENLNKDAEKKSRFLGIFNFMK
 FQKDRKDNZ

f150.nt

TABLE 1. Nucleotide and Amino Acid Sequences

ATGAAAACATTGTTATTATTGGACTTAGTAATTAGGCATTCACTTACTTGAAGATTTAAGCAGGCTTGATTGTC
 AAATTATTATTATAGATACATCTAAAGAGCTTATTGAAGAATATGATGTGATATCTACAGAAAGCTTGTGTTG
 GCAATTCACTAAAATGCTTGTGAAAAGAATAATTCCAGTAGATACAGACGCTGTTGTTATTGATTTGATGATG
 CTTGGCAAAAGTGTCTTACTCACTATTGTAATCTTTAGGTTGAAAGAAATATGCGTTAAGACAGAAAATA
 GAGATGATGCTGAAATCTTAAACTCTTGGGCAACAAAATTATATTCCAAGTAAGATGCTGCAAGAAGATT
 AACTCCATTATTAGTATGCTCAAATCTTCAACTTATAATTATTGGGTATGATATTGTTGCTGAAACTGTT
 ATTCCCAAAGAATATGTTGGTAAAACCTCTTGAAGCCGATCTAGAAGAGAATGTGGGATTACAGTTATGCTG
 TTAGAAATTAAAGTAATTCTAGGTATGAATTGTTGATGGCGATTATTTTTAAAAGATGATAAAATTGTAAT
 TTGTTGAAACAGATAGCATTGAAAATTTCACAAATAATAAAGATTTAATTAAAGATTTAATTTCAGGCTCTAA
 GAGGATGAAAATTAAATAAGATGCTGAGAAAAATCTAGATTAGGATTTCACATTATGAAAATTTC
 AAAAGATCGTAAGGATAATTAG

t150.nt

TGTCAAATTATTATTAGATACATCTAAAGAGCTTATTGAAGAATATGATGTGATATCTACAGAAAGCTTGTG
 TTGAGCAATTCACTAAAATGCTTGTGAAAAGAATATTCCAGTAGATACAGACGCTGTTGTTATTGATTTGATG
 TGATCTGGAAAAGTGTCTTGTAACTCACTATTGTAATCTTTAGGTTGAAAGAAATATGCGTTAAGACAGAA
 AATAGAGATGATGCTGAAATCTTAAACTCTTGGGCAACAAAATTATATTCCAAGTAAGATGCTGCAAGAA
 GATTAACCTCATTATTAGTATCTCAAATCTTCAACTTATAATTATTGGGTATGATATTGTTGCTGAAAC
 TGTATTCCCAAAGAATATGTTGGTAAAACCTCTTGAAGCCGATCTAGAAGAGAATGTGGGATTACAGTTATT
 GCTGTTAGAAAATTAAAGTAATTCTAGGTATGAATTGTTGATGGCGATTATTTTTAAAAGATGATAAAATTG
 TAATTGTTGTAACCAAGATAGCATTGAAAATTTCACAAATAATAAAGATTTAATTAAAGATTTAATTTCAGGCTC
 TAAAGAGGATGAAAATTAAATAAGATGCTGAGAAAAATCTAGATTAGGATTTCACATTATGAAAATT
 TTTCAAAAGATCGTAAGGATAATTAG

f219.aa

MLIARIMNINTLFYGMIIIFALISCNHKNIQYDKRIKKFLDKNKIEYKIDSENDFIANKDINNNEKEEVIIRSRL
 NSYKNSKIREIFGIVKVFDINTPKIKEISDSLMSDSYNNRVFGSWEIIHNAERGINSVYIVKAEEFANDTFLDA
 IDEIASTISIFKKIITNNENIDNNNEENNNTNESNEQPTLKQEKTNSTKESNNELKEDQIEELQEIKAQZ

t219.aa

CNHKNIQYDKRIKKFLDKNKIEYKIDSENDFIANKDINNNEKEEVIIRSRLNSYKNSKIREIFGIVKVFDINTPKI
 KEISDSLMSDSYNNRVFGSWEIIHNAERGINSVYIVKAEEFANDTFLDAIDEIASTISIFKKIITNNENIDNN
 EENNNTNESNEQPTLKQEKTNSTKESNNELKEDQIEELQEIKAQZ

f219.nt

ATGCTAATTGCAAGAATAATGAATATTACATTACTACGGCATGATCATTATCATTGGCACTCATTCTT
 GCAATCATAAGAATATACAGTACGACAAGAGAATTAAAAAATTGGATAAAAACAAAATTGAATATAAAATAGA
 CTCAGAAAATGACTTATAGCATTAAAGATATAACAAATAACGAAAAGAAGAAGTAATCATCAGATCAAGACTA
 AACTCATATAAAAATTCAAAGATAAGAGAAATTGGATTGTTAAAGTATTGATATAAACACACACAAAATAA
 AAGAAATATCTGACTCGTTATGAGCGATAGTTATAAACAGAGTATTGGATCGTGGGAGATTATTCATAATGC
 AGAAAGAGGAATCAACTCTTGGTATATATTGTTAAAGCAGAAGAATTGCAATGATACATTGGCTTGATGCC
 ATTGATGAGATTGCCTCAACAATAAGTATTTCAAAAAAATAAACACACACACACACACACACACACACAC
 AAGAAAATAACAATACAATGAATCAAATGAACAGCCCACCTTAAAGCAAGAAAAACAAATTCAACAAAAGAATC
 TAATAACGAACTTAAAGAAGATCAAATAGAAGAAGACTTCAAGAAATCAAAGCCCACCTTAAAGCAAGAAAAACAAATTCAACAAAAGAATC

t219.nt

TGCAATCATAAGAATATACAGTACGACAAGAGAATTAAAAAATTGGATAAAAACAAAATTGAATATAAAATAG
 ACTCAGAAAATGACTTATAGCATTAAAGATATAACAAATAACGAAAAGAAGAAGTAATCATCAGATCAAGACT
 AACTCATATAAAAATTCAAAGATAAGAGAAATTGGATTGTTAAAGTATTGATATAAACACACACACAAAATA
 AAGAAAATATCTGACTCGCTTATGAGCGATAGTTATAAACAGAGTATTGGATCGTGGGAGATTATTCATAATGC
 AGAAAGAGGAATCAACTCTTGGTATATATTGTTAAAGCAGAAGAATTGCAATGATACATTGGCTTGATGCC

TABLE 1. Nucleotide and Amino Acid Sequences

AATTGATGAGATTGCCTCAACAATAAGTATTTCAAAAAAATAATAACAACCAACAACGAAAACATTGATAATAAT
GAAGAAAATAACAATACAAATGAATCAAATGAACAGCCCACCTAAAGCAAGAAAAACAAATTCAACAAAAGAAT
CTAATAACGAACCTAAAGAAGATCAAATAGAAGAAGAACTTCAAGAAATCAAAGCCCATAAA

f229.aa

MRVDLLPLVELSLYINLSFCKDFSIFNRILEELKCHLILLGPIIKTLYIKHVDFCLSRQDNLKFIITSLSKYIN
LELEEFITLEIIPGYVDFEKFKLDEFCTIRINLNQSFSLERKIVGipeISYKKLNILINNIRKFPFDLNIDMT
VNMPQLQKSHLKRDQLQRIAFIYAZ

t229.aa

CKDFSIFNRILEELKCHLILLGPIIKTLYIKHVDFCLSRQDNLKFIITSLSKYINLELEEFITLEIIPGYVDFEK
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IYAZ

f229.nt

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TTTTAATAGAATTAGAGGAATTAAATGTCATTAACTTGTGGTCATCCAATTATAAAACACTTACAT
TAAGCACGTAGATTTGTTATCTAGGCAAGATAATTAAATTACTTGTCAAGTATATTAAAT
TTGGAGTTATTAGAAGAATTACTTGTAGAAATTATCCGGTTATGTTGAAATTCAAACCTTGGATG
AATTGTTGATTACTAGAATTAACTTAAATGTTGAAATTAGAAAGTTCTTGTAGAAAGATTGTTGGGATACCGA
AATTCTTATAAAATTGAATATTGATTAACAATTAGAAAGTTCTTGTAGAAATTGACATGACT
GTCAATATGCCTTGCAAAAAATCTCATCTCAAGCGAGATTGCAAAGAATTGCTTCATATGCCTGA

t229.nt

TGTAAAGATTAGCATTAAATAGAATTAGAGGAATTAAATGTCATTAACTTGTGGTCATCCAATTAA
AAAAACACTTACATTAAGCACGTAGATTTGTTATCTAGGCAAGATAATTAAATTATTCACCTCTT
GTCCAAGTATATTAAATTGGAGTTAGAAGAATTACTTGTAGAAATTATCCGGTTATGTTGAAATT
TCAAACTTGGATGAATTGTTGATTACTAGAATTAACTTAAATGTTGAAAGTTCTTGTAGAAAGA
TTGTGGGATACCGAAATTCTTATAAAATTGAATATTGATTAACAATTAGAAAGTTCTTGTAGAAAGA
GAATATTGACATGACTGTCAATATGCCTTGCAAAAAATCTCATCTCAAGCGAGATTGCAAAGAATTGCTTC
ATATATGCCTGA

f22.aa

MLKTLTKIITISCLIVGCASLPYTPPKQNLNYLMELLPGANLYAHVNLIKNSIYNSLSPKYKSVLGLISNLYFSY
KKENNDFAALLIMGNFPKDIFWGITHKNRNTESIGNIFTNPWKWLKNSNIYIIPNKARTSIAITQKDITAKDNNMLTT
KYIGEIEKNEFFWIQDPTLLLNPQIVSSKNLIPFSSGTLINSLNQEEYIFKSLIKTNNPILKILSKKLIPTVL
TNMTNLTISSHIKTTIKDQNTVEIEFNIQKSSVESLIEKLASNIQT

t22.aa

CASLPYTPPKQNLNYLMELLPGANLYAHVNLIKNSIYNSLSPKYKSVLGLISNLYFSYKKENNDFAALLIMGNFPK
DIFWGITHKNRNTESIGNIFTNPWKWLKNSNIYIIPNKARTSIAITQKDITAKDNNMLTTKYIGEIEKNEFFWIQD
PTLLLNPQIVSSKNLIPFSSGTLINSLNQEEYIFKSLIKTNNPILKILSKKLIPTVLTNMTNLTISSHIKTTIK
DQNTVEIEFNIQKSSVESLIEKLASNIQT

f22.nt

ATGTTAAAAACATTAACAAAAATAATTACCATTCATGCCTCATAGTGGATGCGCAAGCCTGCCTACACTCCTC
AAAAACAAATCTAAATTACTTAATGAACTTTACCTGGCGAAATTATACGCCATGTAATTAAATTAAAAAA
CAGGTCTATTATAACTCTTAAGCCCTAAATATAAATCAGTCTTGGCTTATAAGCAATTATACTTAGCTAT
AAAAAAGAAAATAACGATTTGCTACTAATAATGGTAATTCCAAAAGATATTCTGGGAATTCAAAAAA

TABLE 1. Nucleotide and Amino Acid Sequences

ATAGAAATACAGAATCAATAGGCAATATATTACAAATCCAAAATGGAAACTTAAAAATTCAAATATATACATTAT
 TCCAAACAAAGCTAGAACTAGCATTGCAATAACCCAAAAGATATAACCGCAAAAGACAATAATGCTAACAA
 AAATATATTGGGAAATAGAAAAATGAAATGTTTTGGATTCAAGATCCAACATTATTGCTCCCAACCAAA
 TAGTAAGCAGCAAAATTTAATCCCTTAGCAGTGAACCTTGTCTATAAACAGCTTAAATCAAGAAGAATATAT
 TTTTAAATCCTTAATCAAAACAATAATCCACCAATACTAAAATATTGTCAAAAAGTTAATCCAACCGTCTG
 ACAAAACATGACAAACCTCACAAATATCAAGCCACATAAAGACCACAATAAAGACCAAAATACGTTGAAATAGAAT
 TTAATATTCAAAAATCTAGTGTGAAAGCCTTATAGAAAAACTAGCTCAAATATTCAAACCTAA

t22.nt

TGCGCAAGCCTGCCTACACTCCTCCAAAACAAAATCTAAATTACTTAATGGAACCTTACCTGGCGCAAATTAT
 ACGCCCATGTAATTTAATTAAAAACAGGTCTATTAACTCTTAAGCCCTAAATATAATCAGTTCTGGGCT
 TATAAGCAATTATACATTAGCTATAAAAAGAAAATACGATTTGCTCTACTAATAATGGGTAAATTCCCAAA
 GATATTCTGGGAATTCAATAAAATAGAAATACAGAATCAATAGGCAATATATTACAAATCCAAAATGGAAAC
 TTAAAATTCAAATATACATATTCCAAACAAAGCTAGAACTAGCATTGCAATAACCCAAAAGATATAACGC
 AAAAGACAATAATATGCTAACAAACAAATATATTGGGAAATAGAAAAAAATGAAATGTTTTGGATTCAAGAT
 CCAACATTATTGCTCCAAACCAAAATAGTAAGCAGCAAAATTAAATTCCCTTACAGTGGAACTTGTCTATAA
 ACAGCTTAAATCAAGAAGAATATATTAAATCCTTAATCAAAACAAATAATCCACCAATACTAAAATATTGTC
 AAAAGTTAATTCAACCGTCTGACAACATGACAAACCTCACAAATATCAAGCCACATAAAGACCCACAATAAAA
 GACCAAAATACGGTTGAAATAGAATTAAATTCAAAAATCTAGTGTGAAAGCCTTATAGAAAAACTAGCTCAA
 ATATTCAAACCTAA

f32.aa

MNTKTLYLISLILLACNKNKIPLIQKLDLPKSSILGFSNKMGIICKDYAFLSKSTKNSELDYDYAILLRKDEVV
 KIEKTLKTERYGIEGNWILVNYKGTKRYIFSKDINIVNNLIDHSK

t32.aa

CNKNNKIPLIQKLDLPKSSILGFSNKMGIICKDYAFLSKSTKNSELDYDYAILLRKDEVVKIEKTLKTERYIE
 GNWILVNYKGTKRYIFSKDINIVNNLIDHSK

f32.nt

ATGAATACAAAACATTATATTAAATCCTTAATTCTTTAGCTTGCAATAAAAATAACAAAATTCTCTCATTC
 AAAATTAGATTGCCCCAAAGCAGCATTCTGGCTTAGCAATAAAATGGGCATAATAATAAAAGATTATGCTTT
 TCTTAGTAAAGCACTAAGAAAAATAGCGAATTGGATTATGATTACGCAATTCTACTCAGAAAAGACGAAGTCGA
 AAAATTGAAAAACACTAGAAAAACAGAGCGCTATGGAATTGAAGGAAATTGGATCCTAGTCATTACAAGGGAA
 CTTAAAGATACATCTTAGCAAAAGACATCAATATAGTCACAAATTAAATTGATCATTCTAAATAG

t32.nt

TGCAATAAAAATAACAAAATTCTCTCATTCAAAATTAGATTTGCCAAAAGCAGCATTCTGGCTTAGCAATA
 AAATGGGCATAATAATAAAAGATTATGCTTTCTTAGTAAAGCACTAAGAAAAATAGCGAATTGGATTATGATTA
 CGCAATTCTACTCAGAAAAGACGAAGTCGAAAAATTGAAAAACACTAGAAAAACAGAGCGCTATGGAATTGAA
 GGAAATTGGATCCTAGTCATTACAAGGAACTAAAAGATACATCTTAGCAAAAGACATCAATATAGTCACAAATT
 TAATAATTGATCATTCTAAATAG

f186.aa

MKKLIIIFTLFLSQACNLSTMHKIDTKEDMKILYSEIAELRKKLNHNLEIDDTLEKVAKEYAIKLG
 ENRTITHLFGTTPMQRHKYDQSFNLTREILASGIELNRVVAWLNSPSHKEALINTDTDKIGGYRLKTTDNIDIFV
 VULFGKRKYKN

t186.aa

TABLE 1. Nucleotide and Amino Acid Sequences

CNLSTMHKIDTKEDMKILYSEIAELRKKLNLNHLEIDDTLEKVAKEYAIKLGENTHTLFGTT
 PMQRIHKYDQSFNLTREILASGIELNRVVAWLNSPSHKEALINTDTDKIGGYRLKTTDNIDIFVVLFGKRKYKN

f186.nt

ATGAAAAAAATTGATTATAATTTTACACTGTTTATCTCAAGCATGCAATTAAAGTACAATGCATAAAATAGATA
 CAAAAGAAGATATGAAAATTCTATATTCAAGAAATTGCTGAATTGAGAAAAAAATTAAATCTAAACCATCTAGAAAT
 AGATGATACCCCTGAAAAAGTTGCAAAAGAATATGCCATTAAACTGGGAGAAAATAGAACAAACTCACACCCCTT
 TTTGGCACACCCCAATGCAAAGAATACATAAATACGATCAATCCTTAATTAAACAAGAGAAATACTGGCATCAG
 GAATTGAACCTAACAGAGTAGTTAATGCATGGCTTAATAGTCCAAGCCACAAAGAACAGCTCTTATTAAACAGATAC
 CGATAAAATAGGTGGCTATAGATTAACAGACTGACAATATAGATATTGTAGTTCTTTGGAAAAAGAAAA
 TATAAGAATTGA

t186.nt

TGCAATTAAAGTACAATGCATAAAATAGATACAAAAGAAGATATGAAAATTCTATATTCAAGAAATTGCTGAATTGA
 GAAAAAAATTAAATCTAAACCATCTAGAAATAGATGATACCCCTGAAAAAGTTGCAAAAGAACATGCCATTAAACT
 GGGAGAAAATAGAACAAACTCACACCCCTTTGGCACACCCCAATGCAAAGAACATACATAAATACGATCAATCC
 TTTAATTAAACAAGAGAAATACTGGCATCAGGAATTGAACTTAACAGAGTAGTTAATGCATGGCTTAATAGTCCAA
 GCCACAAAGAAGCTTTATTAAACAGATACCGATAAAATAGGTGGCTATAGATTAACAGACTGACAATATAGA
 TATATTGTAGTTCTTTGGAAAAAGAAAATATAAGAATTGA

f216.aa

MIRVLLGSLAVSFLSICMVFLNYDNLFSSKKVYFHSSKGFBANLRYLRDEQNLKDNLVDFLLGSNEGFSFG
 FLLSDSRFLYSFLKNGVYYVNLREFYDSFNNGDYNESNESFDVKVNLFAMSLIKTMRFNYPGKIKKIVILVEGCI
 LKEQS

t216.aa

CMVFLNYDNLFSSKKVYFHSSKGFBANLRYLRDEQNLKDNLVDFLLGSNEGFSFGFLSDSRFLYSFLKNGV
 YYVNLREFYDSFNNGDYNESNESFDVKVNLFAMSLIKTMRFNYPGKIKKIVILVEGCI
 LKEQS

f216.nt

ATGATTAGGGTGCTTTGGGTCTTGGCAGTAAGCTTTGTTCTATTGTATGGTTTTAAATTATGATA
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 ACAAAATTGAAAGATAATTAGATCTTAGTAAAAGATTTCTTTAGGAAGCAATGAAGGGTTCTTTGGG
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 TTTATGATTCTTAATAATGGTATTATAATGAATCTAATGAATCTTGATGTTAAGGTCAATCTTTGCTAT
 GTCTTTAAACATGCGCTTAACATCCTGTAAGATAAAAGATTGTTATTCTGTTAAGGGTGTATC
 TTAAAGGAGCAAAGTTGA

t216.nt

TGTATGGTTTTAAATTATGATAATCTTTCAAAAAGGTTTTATTTCAATTCTAGCAAGGGATTGTTG
 CTAATTAAAGATATTAAAGAGATGAACAAAATTGAAAGATAATTAGATCTTTAGTAAAAGATTTCTTTAGG
 AAGCAATGAAGGGTTCTTTGGTTTATTAAAGTGAATCAAGATTATATTCTTTAAAGAACATGGAGTT
 TATTATGAAATCTTCAAGAGAATTATGATTCTTTAAATGGTATTATAATGAATCTAATGAATCTTTG
 ATGTTAAGGTCAATCTTTGCTATGTTAATAACATGCGCTTAACATCCTGTAAGATAAAAGAT
 TGTTATTCTGTTAAGGGTGTATCTTAAAGGAGCAAAGTTGA

f328.aa

MAIKYARENINPFLGICLGLQLAVIEFARNVCGILDADTEENLARDKPLKSPVHLLPEQKGIKDKGATMRLGGP
 VILKKNTIAFKLYGQDRIIERFRHRYEVNNDYIDLFAKNGLIVSGFSSDFKMAKLIEIPENKFFVACQFHP
 IENPAKLFGLIKACI

TABLE 1. Nucleotide and Amino Acid Sequences

t328.aa

CLGLQLAVIEFARNVCGILDADTEENLARDKPLKSPVIHLLPEQKGIKDKGATMRLGGYPVILKKNTIAFKLYGQD
 RIERFRHRYEVNNDYIDLFAKNGLIVSGFSSDFKMAKLIEIPENKFFVACQFHPELITRIENPAKLF
 LGLIKACI

f328.nt

ATGGCTTAAATATGCTCGTGAGAATAATTCCCTTCTTGGAAATTGTCTTGGTTGCAGCTTGCTGTAATAG
 AATTGCTCGTAATGTTGGAAACTTGATGCTGATAACGGAGGAAAATTAGCAAGAGACAAGCCCTAAAAAG
 TCCTGTTATCCATTACTTCCTGAGCAAAGGGAAATAAGATAAGGGCCTACAATGAGGCTGGTGGATATCCT
 GTGATTCTAAAAAGAATAACAATAGCTTTAAACTTATGGCAAGATGGATAATTGAAAGATTAGACATAGGT
 ATGAAGTCATAATGATTATAGATTTAGCTGAAAAATGGCTTATAGTATCTGGATTTCAAGTGA
 ATGGAAAATTAGAAATTCTGAAATAATTTCGTAGCTTGCAGTTCCATCCAGAACTTACAAGA
 ATAGAAAATCCAGCCAAGCTTTCTAGGATTAATTAAAGCTTGTATTGA

t328.nt

TGTCTGGTTGCAGCTTGCTGTAATAGAATTGCTCGTAATGTTGTGAAACTTGATGCTGATAACGGAGGAAA
 ATTAGCAAGAGACAAGCCCTAAAAAGTCCTGTTATCCATTACTTCCTGAGCAAAGGGAAATTAAAGATAAGGG
 CGCTACAATGAGGCTGGTGGATATCCTGTGATTCTAAAAAGAATAACAATAGCTTTAAACTTATGGCAAGAT
 CGGATAATTGAAAGATTAGACATAGGTATGAAGTCATAATGATTATAGATTTAGCTGAAAAATGGCTTA
 TAGTATCTGGATTTCAAGTGAATTGGCAAATTAGAAATTCCGTGAAATAATTTCGTAGCTTGC
 CCAGTTTCATCCAGAACTTACAAGAATAGAAATCCAGCCAAGCTTTCTAGGATTAATTAAAGCTGTATT
 TGA

f352.aa

MNKTNRSLTYFIILSCISLFGANNNTISYSSIEIPLEDLSEEFKSSGNKSDQINTSKHLNKNIVSYEDPKKGKDL
 KL PENIRDKKLPQKRM DENDLKS VIEN YEN KI NIE KLLK TKNQ KTS ENEN KI EIE KKK YE I LTNKL KNE IV
 EIKKLLNKKIKPKEDENYEKINIEENIEETDDDFEDNYEYDEIEXTNEDNYPSEGIINNLKENLNENEKYAIN
 EKKIDELEDRINENENTILDQLRELRNFKKDKNSDKNLEEEENLSSIGRIINDLKRKISANEAINKENQKKIRTD
 KHKLKELEDKIKENEETILKLQKELNNFKKKKEIYQKPLNEETFTPSITSKNDDEENKKLKEYLKPIEKKESRDL
 EENTKSTPKTTMIKTADFQIYDPIYLNYYKFKEKGDFQFAFKKENTYYIEIDPTNNLNEALKNHEIISKYKFEKYFI
 NPILKNKEEFFRNLLIEVKNIHELGIMYKNLKEFKQIKIIK

t352.aa

CISLFGANNNTISYSSIEIPLEDLSEEFKSSGNKSDQINTSKHLNKNIVSYEDPKKGKDLKL PENIRDKKLPQKRM
 DENDLKS VIEN YEN KI NIE KLLK TKNQ KTS ENEN KI EIE KKK YE I LTNKL KNE IV EIKKLLNKKIKPKED
 NYE KINIEENIEETDDDFEDNYEYDEIEXTNEDNYPSEGIINNLKENLNENEKYAIN EKKIDELEDRINENEN
 TILDQLRELRNFKKDKNSDKNLEEEENLSSIGRIINDLKRKISANEAINKENQKKIRTDKHKLKELEDKIKENE
 TILKLQKELNNFKKKKEIYQKPLNEETFTPSITSKNDDEENKKLKEYLKPIEKKESRDL EENTKSTPKTTMIKTA
 DFQIYDPIYLNYYKFKEKGDFQFAFKKENTYYIEIDPTNNLNEALKNHEIISKYKFEKYFINPILKNKEEFFRNLL
 VKNIHELGIMYKNLKEFKQIKIIK

f352.nt

ATGAATAAAACAAAAATCGAAGCCTACGTATTTATAATACCTTACATGATATCATTATTGGGGCTAATAATA
 ATACAATAAGCTACTCTAGCATTGAAATTCTCTAGAAGACTTAAGTGAAGAATTAAAAGTCTGGGAATAAAAG
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 AAATTGCCAGAAAATATAAGAGACAAAAACTACCCAAAAAGAATGGACGAAATGATCTAAATCTGAATTG
 AAAATTATGAAAATAAAATTTAAACATAGAAAAGCTTTAAAACCAAAAATCAAAAACATCGGAAATGAAA
 TAAAAAAATAGAATCAATCGAAAAAAGCAAAAAATGAAATTAAACCAATAATTAAAACGAAATAGTA
 GAAATAAAAAGCTCCTAACAAAAATCAAGCTAAAGAAGATGAAAATTACGAAAAATAATATTGAAAACA

TABLE 1. Nucleotide and Amino Acid Sequences

TTGAAGAAGAAACTGATGATGATTTGAAGACAATTATGAATATAATGATGAAATTGAGAACAAATGAGGACAAT
 TACCCCTCTAATGAAGGAATAATAACAATCTAAAAGAAAATCTTAATGAAAACGAAAATATTATGCTATTAATG
 AAAAAAAATCGATGAACCTGAAGACAGAACGAGAATGAAAACACTATTTAGACTTGCAAGAGAAATTAAG
 GAATTTAAAAAAAGATAACTCAGATAAAACTTAGAAGAATTGAGGAAAATTATCTCAATAGGAAGAATA
 ATTAATGATCTAAAAGAAAATCAGCGAAATGAAGCAATAACAAAGAAAATCAAAAAAAATAAGAACTGATA
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 CAATTTAAAAAAAGAAATTATCAAAACCTTAAATGAAGAACTTCACTCCAAGCATTACAAGTAAAAT
 GACGACTTAGAAGAAAATAAGAAATTAAAAGGAATATTTAAGCCATAGAAAAAGAAAGCCGAGATCTAG
 AAGAAAATACTAAAAGCACCCAAAACAATGATAAAACAGCAGATTCTAACTACCCGTACATATATCT
 TAATAATTATAATTAAAGAAAAGGGAGATCAATTGCATTAAAAAGAAAACACATACTATATTGAAATAGAT
 CCCACTAACAAATTAAATGAGGCTTAAAAATCATGAAATACTCAAATATAAATTGAAAATATTCATTA
 ACCCTATTCTAAAAATAAGAAGAATTTTAGAAACTTAATAGAAGTCAAAATATCCACGAACCTAGGAATTAT
 GTATAAAAATCTAAAGCCTGAATTAGCAAATAAAATAATTAAATA

t352.nt

TGTATATCATTATGGGGCTAATAATAACAATAAGCTACTCTAGCATTGAAATTCCCTAGAAGACTTAAGTG
 AAGAATTAAAGTTCTGGATAAAAGCGATCAAATAACCTCAAAACATTAAACAAAATAGTTCTTA
 TGAAGACCCAAAAGGGTAAAGATCTAAAATTGCCAGAAAATATAAGAGACAAAACACTACCCAAAAGAATG
 GACGAAAATGATCTAAATCTGTATTGAAAATTATGAAAATAAAATTAAAACATAGAAAAGCTTTAAAACCA
 AAAATCAAAAACATCGGAAAATGAAAATAAAATAGAATCAATGAAAAAGCAAAAATATGAAATT
 AACCAATAATTAAAAGAAATAGTAGAAAATAAAAGCTCTTAACAAAAAAATCAAGCCTAAAGAAGATGAA
 ATTACGAAAATAATATTGAAAACATTGAAGAAGAACTGATGATGATTTGAAGACAATTATGAAATATAATG
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 CTATTTAGACTGCAAAGAGAATTAGGAATTAAAAGATAACTCAGATAAAACTTAGAAGAAATTGAA
 GAAAAATTATCTCAATAGGAAGAATAATTATGATCTAAAAGAAAATCAGCGAAATGAAGCAATAACAAA
 GAAAATCAAAAATAAGAAACTGATAAACACAAACTCAAAGAATTAGAAGATAAAATAAGAAAATGAAGAGA
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 TTTCACTCCAAGCATTACAAGTAAAATGACGACTTAGAAGAAAATAAGAAATTAAAAGGAATTATTAAGCC
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 AGAAAACACATACTATATTGAAATAGATCCACTAACATTAAATGAGGCTTAAAAATCATGAAATTATCTCA
 AAATATAAATTGAAAATATTCATTAACCCATTCTAAAGAAGAATTTTAGAACTTAATAGAAG
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 A

f867.aa

MNTKGKVVGVNGNLVTIEVEGSVSMNEVLFVKTAGRLKAEVIRIRGNEVDAQVFELTKGISVGLVEFTDKLLTV
 ELGPGLLTQVYDGLQNPLPELAIQCGFFLERGVYLRPLNKKWNFKKTSKVGDIVIAGDFLGVIETVHHQIMI
 PFYKRD SYKIVEIVSDGDY SIDEQIAVIEDDSGMRHNIITMSFHWPVKVPITNYKERLIPSEPM LTQTRIIDTFFPV
 AKGGTFCIPGPFGAGKTVLQQVTSRNADVDVIIAACGERAGEVVELKEFPELMDPKTGKSLMDRTCIICNTSSM
 PVAAREASVYTAITIGEYYRQMLDILLADSTS RWAQAMREMSGRLEEI PGEEAF PAYLESVIASFYERAGIVVL
 NNGDIGSVTVGGSVSPAGGNFEEPVQTQATLKVVGAFHGLTRERSDARKFPAISPLESWSKYKGVIDQKKTEYARSF
 LVKGNEINQMMKVVGEEG ISNDDFLIYLKSELLDSCYLQQNSFDSIDA AVSSERQNYMFDIVYNILKTNFEFSDKL
 QARDFINELRQNLLDMNLSSFKDHFKNLEHALGELINFKKVI

t867.aa

GRNLKAEVIRIRGNEVDAQVFELTKGISVGLVEFTDKLLTVELGPGLLTQVYDGLQNPLPELAIQCGFFLERGVY
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 RHNIITMSFHWPVKVPITNYKERLIPSEPM LTQTRIIDTFFPVAKGGTFCIPGPFGAGKTVLQQVTSRNADVDVII
 AACGERAGEVVELKEFPELMDPKTGKSLMDRTCIICNTSSMPVAAREASVYTAITIGEYYRQMLDILLADSTS
 RWAQAMREMSGRLEEI PGEEAF PAYLESVIASFYERAGIVVLNNGDIGSVTVGGSVSPAGGNFEEPVQTQATLKVG
 AFHGLTRERSDARKFPAISPLESWSKYKGVIDQKKTEYARSFLVKGNEINQMMKVVGEEG ISNDDFLIYLKSELLD

TABLE 1. Nucleotide and Amino Acid Sequences

SCYLQQNSFDSIDA AVSSERQNYMF DIVYNILKTNF EFS DKLQARDF INELRQNLLDMNLSSFKDHKF NKL EHALG
ELINFKKVI

f867.nt

ATGAATACAAAAGGAAAAGCTGGAGTTAATGGAACTTAGTTACTATTGAGGTAGAAGGTTCA GTTTCTATGA
ATGAAGTTTATTTGTAAGACTGCTGGTAGGAATTAAAAGCAGAAGTAATCGTATTAGGGCAATGAAGTTGA
TGCACAGGTTTGTAAAGACAAAAGGGATATCTGTTGGAGACCTAGTTGAATTACAGACAAACTTTAACAGTT
GAACTCGGACCAGGGCTTTAACGATATGATGGGCTCAAAATCCTTGCGCTGAATTGCGTATTCAATGTG
GATTTTTTAGAAAGGGAGTATATTAAAGGCCCTGAATAAAGATAAAAAGTGAATTAAAAACCTCCAA
AGTTGGAGATATCGTTATTGAGGAGATTTTAGGTTGTAAATTGAGGAACGTGTCACCACAAATAATGATT
CCATTTATAAAAGGGATTCTTATAAAATTGTGGAGATTGTAAAGTGAATGGCAGACTATTGATGAGCAAATTG
CTGTAATTGAAGATGATTCTGGTATGAGGCATAATATTACAATGTCTTTCAATTGGCCTGTTAAAGTCCATTAC
TAATTATAAGGAACGCCTTATTCCCTAGTGAACCTATGTTGACTCAAACATAGAATTATAGATACATTTCAGTT
GCCAAAGGTGGAACTTTGCATTCCGGCTTTGGACCGAGGAAAACGGTTCTCAGCAGGTTACAAGTCGAA
ATGCTGATGTTGATGTAGTATTGAGCTTGTGGTGGAGCAGGAGAGAAGTGGTAGAAACTCTTAAAGAATT
TCCCAGATTAAATGGATCCAAAACCGGAAATCTTAATGGACAGGACTTGTATTGATCTTAAACATCTCAATG
CCAGTTGCACTAGAGAAGCTCTGTTACTGCTATTACTATTGGTGGAGTATTACAGGAAATGGCCTTGATA
TTCTCTTGGCAGATTCAACTCAAGATGGGCTCAAGCAATGAGAGAAATGTCTGGACGCCCTGAGGAAATTCC
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CTCAAGCAACTTTAAAGTGTAGGAGCATTTCACGGGCTTACAAGAGAAAGGTCTGATGCTAGGAAATTCCAGC
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TTTATTAAATCCGAGCTACTGATTGCTATTGAGCAAAATTCAATTGATCTTAAACTTGAAGTCTGATAAAACTT
TTCAGAGCGTCAAAATTATATGTTGATATAGTTATAACATTCTAAACTTGAGTTCTGATAAAACTT
CAAGCAAGAGATTTTATAAATGAGTTAAGGCAAAATTCTTACAGATGAATCTTCTCTTTAAGGATCATAAGT
TTAATAATTGGAGCATGCTTGGGTGAATTGATAAATTAAAAGGTAAATTAG

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GGTAGGAATTAAAAGCAGAAGTAATCGTATTAGGGCAATGAAGTTGATGCACAGGTTTGAATTGACAAAAG
GGAT
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GGGCTCAAGCAATGAGAGAAATGTCTGGACGCCCTGAGGAAATTCTGGAGGAGGCTTCCGGCATATCTG
GTCTGTTATTGCTCCTTTATGAAAGGGCAGGTATTGAGTTCTTAATAATGGGATATTGGATCTGTAACAGT
GGTGGCTCTGTAAGTCCTGCTGGTAATTGAGAGGCCAGTTACTCAAGCAACTTTAAAGTTGAGGAGCAT
TTCACGGGCTTACAAGAGAAAGGTCTGATGCTAGGAAATTCCAGCTATTGCTCTGTAATTGAGGAGATA
TAAAGGCGTTATTGATCAAAAAAAAGACTGAATATGCAAGATCTTTTGAGGAAAGGTAAATGAAATTAAATCA
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GCTATTGAGCAAATTCAATTGATTCTATTGATGCTGCTGTTAGTTGAGCGTCAAATTATGTTGAT
AGTTTATAACATTCTAAACTTGTAGTTCTGATAAAACTTCAAGCAAGAGATTAAATGAGTTAAGG
CAAATCTTACAGATGAATCTTCTCTTTAAGGATCATAAGTTAATAATTGGAGCATGCTTGGGTGAAT
TGATAAATTAAAAGGTAAATTAG

TABLE 1. Nucleotide and Amino Acid Sequences

f868.aa

MKRVYSKIESIAGNVITVTAQGIKYGELAIVKAKDTSSLAEVIKLDREKVSLQVYGGTRGVSTSDEIKFLGHSMQVSFSDNLLGRIFDGSGNPRDGPSLDDNLIEIGGPSANPTKRIVPRNMIRTGLPMIDVFTLVEQKLPPIFSVSGEPYNELLIRIALQAEVDLILIGMGLKHDDYLTFKDSLEKGALSRAlFFVHTANDSVESLTVPDISLSVAEKFALKGKVLVLLTDMTNFADAMKEISITMEQVPSNRGYPGDLYSQLAYRYEKAIIDFEGAGSITILAVTTMPGDDVTHPVDNTGYITEQQYVLKGGRIEPFGSLSRLKQMVNSRTRDDHRTIMDSMIKLYASSKESVEKKAMGFNMTKWDEKLKYSNMFESKMMDSLNVNIPLEEALDLGWSILASCFSPKETGIKTDLIEKYWPKKETY

t868.aa

QGIKYGELAIVKAKDTSSLAEVIKLDREKVSLQVYGGTRGVSTSDEIKFLGHSMQVSFSDNLLGRIFDGSGNPRDGPSLDDNLIEIGGPSANPTKRIVPRNMIRTGLPMIDVFTLVEQKLPPIFSVSGEPYNELLIRIALQAEVDLILIGMGLKHDDYLTFKDSLEKGALSRAlFFVHTANDSVESLTVPDISLSVAEKFALKGKVLVLLTDMTNFADAMKEISITMEQVPSNRGYPGDLYSQLAYRYEKAIIDFEGAGSITILAVTTMPGDDVTHPVPDNTGYITEQQYVLKGGRIEPFGSLSRLKQMVNSRTRDDHRTIMDSMIKLYASSKESVEKKAMGFNMTKWDEKLKYSNMFESKMMDSLNVNIPLEEA

f868.nt

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AGAGACTTATTGA

t868.nt

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AAAATTGCTCAAGTATGCAATATGTTGAAAGTAAGATGATGGATTGCTGTTAATATTCTTTAGAAGAGGCT

TABLE 1. Nucleotide and Amino Acid Sequences

TTAGATTTAGGTTGGAGCATTCTGCTAGTTAGCCAAAAGAAACGGGAATAAAACAGATCTTATTGAA
AATATTGGCTAAAAAAGAGACTTATTGA

f872.aa

MRSAVLFFFALPFSISLYSSSNKNFPYWILLEKGRQFLYSKSEFSKSNLTHAINYLQEALLRKGVYPEASYYLSVA
YMSGNAILEKLNLYKSFEDRYYLLDESFEKKILFLAKMAELENNYVDTIDYLNDILNKFSTKKDYYSYHDYSQG
ENSMSNNELNASFYLTSLKQVRGAFGIDFTFNLYRFKNYNVIDTHQLLSKVYLHLKAYELSITHGLIAAVGILTR
MYDYVCYYEPVYQFKNLRSFVQKINKYKAIAKNAFESTDFWEIVYNVAAATYAYSNGNYKFRaidTWKLVVDLAPRF
SPYIAKSRSQIKNSVYLKNN

t872.aa

SNKNFPYWILLEKGRQFLYSKSEFSKSNLTHAINYLQEALLRKGVYPEASYYLSVAYMSGNAILEKLNLYKSFED
RYYLLDESFEKKILFLAKMAELENNYVDTIDYLNDILNKFSTKKDYYSYHDYSQGENSMSNNELNASFYLTSLK
QVRGAFGIDFTFNLYRFKNYNVIDTHQLLSKVYLHLKAYELSITHGLIAAVGILTRMYDYVCYYEPVYQFKNLRSF
VQKINKYKAIAKNAFESTDFWEIVYNVAAATYAYSNGNYKFRaidTWKLVVDLAPRFSPYIAKSRSQIKNSVYLKNN

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t872.nt

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TAA

f874.aa

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VVITAGLNQKPGETRLDLVDKNSKIFKDIITNVVSSGFDGIFVVASNPVDIMTYVTMKYSKFPPIHKVIGTGTILD
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TABLE 1. Nucleotide and Amino Acid Sequences

GATYYAIGLGIKNIVNAIIGDQNVLPISSYINGQYGGLIKDIYIGAPAIVCKEGVKEVLNFKISPKELDKFNSSA
NQLKSYIDKMEF

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NSKIFKDIITNVVSSGFDGIFVVASNPVDIMTYVTMYSKFPYKVI GTG TILDTSRLRYFLSDHFNVNTQNIHSY
IMGEHDSSFATWDETAKIAMKPLSEYLAEGKITELELDEIHKKVVNAAYEVIKLKGATYYAIGLGIKNIVNAIIGD
QNVLPISSYINGQYGGLIKDIYIGAPAIVCKEGVKEVLNFKISPKELDKFNSSANQLKSYIDKMEF

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f886.aa

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KITCYDTKDTKRKEEYDNLNNKIQEIEYDSKGKTLETANYVYENENLISKNLKTIQKPKLIYYSKDDNGKLLK
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EYHNDNEYEEEIYNNKKPALRVKHKNKGKVTEEKPIGTN

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SYFASDVFFNKYQKLNEPKTGFYIEVYSVDDTEKLYLYKENNLKYKTIQIIDENTKKITCYDTKDTKRKEEYD
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LRVKHKNKGKVTEEKPIGTN

TABLE 1. Nucleotide and Amino Acid Sequences

f886.nt

ATGAAAAAAAACAATTAATACCTCTTCTATTTATGCCACAAATTATTCAGCAAAAGCTATTTGCATCTGATG
 TATTTTCAATAAATACCAAAATTAAATGAAAACCAAAACGGGTTTATATTGAGTATTATCTGTGATGA
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 TPIEIEKPNQDIILLKSKGYKDKFKLINKEEDQVEIEMIKTNKNRLIDTRDKFYVNLAVFTLSTIGAIFAGTLLN
 NSEVLYKITGNHFINKRLTAEDVYMAKAEQMTATFLFGVGITLTIGSFISLITHLVEYIKEANMGE

f888.nt

TABLE 1. Nucleotide and Amino Acid Sequences

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 AATCAAGATATAATAAAACAGAACTGAAATTAGCAAATTAAAAAGAAATGGATAAAAAAAACTTCAAAACA
 TAATAACCGCAAAGAAAAGCATAACACCAAAACCAAATTGATGAGCTAAAAAAATATTCAAATATAACAA
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 TCTATCACGATAATATTGACATTCTAAACTCCAAACAAAGAAATTGAAATACAATCACAAGTCAAATT
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 TTGAAATTGAAAACCAGAAAATCAAGATATCATCTGCTTAAACTAAAGGATATAAGATAATTCAAGTTAAT
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 AATAACTCATTAGTAGAATATATAAGAAGCAAATATGGGAGAATAG

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 LLIIFLDPTNSIFTLIFLILSSLAFMISKEIMYFYPFTVLSYLLFLIIISNFNKNYNKIYLKEINFLLMTKIKHLLF
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 LLLFTLSSLIPFAYSSYMLNSYENINYLYSKKLNYFDYLNPNNIYIMLGYNKDMNPNIIGYLSHILYQNELKYNITAK
 YGKIPDKIENYFEIKNDKIEIHPKTVYEVDKSFIDEILKKDLASLFLKNKNPILYKENKNNINTDKKNYKILFF
 FSLPFFVLLFLFKAIRFTILLNIN
 EKTYKKYIQC

t893.aa

CDAAQFGDYKPLYFENENDLKTANEYINSLGYKTISEYTTKIDILDFFENKEITINEINKLNNDLRKSIFLKKL
 NLNIEHKKLLYVENRFKSINFKNLKKELNINADIHSLDYKTKINFISIIFLIIIIIFLDPTNSIFTLIFLII

TABLE 1. Nucleotide and Amino Acid Sequences

SSLAFMISKEIMYFPFTVLSYLLFLIISNFNKNYNKIELKEINFLLMTKIKHLLFLFTFTALYFITITFFTTN
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LLNINEKTYKKYIQC

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AAATCCAATCTAATATATAAAGAAAACAAGAATAATATCAACACAGATAAAAAAAATTACAAAATACTTTCTT
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t893.nt

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TTACAAAATACTTTCTTTCTTGTGCCCCCTTGTATTACTATTCTTAAAGCAATAAGATTACAATT
CTTTAAACATAATGAAAAACCTATAAAATATTCAAGGATAA

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MIRALLTNDLFLSCLVSGISAQVIKYGIQTVKTRKLKLTPVHLLKKIFLETGGMPSSHSSVTALSTSIALTEGID
TNFIIIALAFALITIRDGSFGVRYMSGVQAEYLNALSEKLKKEIKIDTTKIKVVKGHKKVEVTGIIIGIVSAYIVCY
F

TABLE 1. Nucleotide and Amino Acid Sequences

t895.aa

AQVIKYGIQTVKTRKLKLTPVHLLKKIFLETGGMPSHSSTVTALSTSIALTEGIDTNFIIALAFALITIRDSFGV
RYMSGVQAEYLNALSEKLKEIKIDTTKIKVVKGKKKEVLTGIIIGIVSAYIVCYF

f895.nt

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ACAAATTTATAATAGCTCTGCATTGCCCTTATTACAATAAGAGATTCTTCGGCGTAAGATATATGCTGGAG
TTCAAGCAGAATATTAAATGCATTATCAGAAAAATTAAAAAGAAATAAAATTGACACAACAAAATAAAAGT
GGTCAAGGGCACAAGAAAGAGGTTCTAACGGGCATAATAATAGGAATAGTCTCGGTATATTGTGTGCTAT
TTTAG

t895.nt

GCTCAAGTGATTAATATGGTATCCAAACTGTAAAACAAGAAAGTAAAACACTCCAGTACATCTTAAAAA
AAATTCTAGAAACAGGAGGCATGCCAAGTAGTCATTCAACGGTCACCGCTCTTCAACCTCAATCGCACT
AACTGAAGGAATAGATACAAATTATAATAGCTCTGCATTGCCCTTATTACAATAAGAGATTCTTCGGCGTA
AGATATATGCTGGAGTTCAAGCAGAATATTAAATGCATTATCAGAAAAATTAAAAAGAAATAAAATTGACA
CAACAAAATAAAAGGGTCAAGGGCACAAGAAAGAGGTTCTAACGGGCATAATAATAGGAATAGTCTCTGC
GTATATTGTGTGCTATTAG

f605.aa

MYIGAAGKSFIIIDSFLNCFLFIGFSRSDSLMSLSNSRFEYPYDASCEFLVNIVKYVCGSKYSPMRPTLII
SKLPVFLLVRTGQFSLVSIRLIFRIFFHWZ

t605.aa

CFLFIGSFRSDSLMSLSNSRFEYPYDASCEFLVNIVKYVCGSKYSPMRPTLIIISKLPVFLLVRTGQFSLVSIR
LIFRIFFHWZ

f605.nt

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TGAATTCTCTGTGAATATAGTAAAGTATGTGTGGATCTAAATATCCCCAATCGGTCAAACCTTTATT
TCAAAATTGCCAGTATTCTGCTGTTGGTAAGAACAGGCCAATTCTGGTAAGCATAAGATTGATATTAGAA
TTTTTTCCATTGGTTTG

t605.nt

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ATGATGCAAGTTGTGAATTCTCTGTGAATATAGTAAAGTATGTGTGGATCTAAATATCCCCAATCGGTCC
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TTGATATTAGAATTCTCCATTGGTTTG

f606.aa

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RAIDEKTLITLESPKPYFIDMLVHQSFIPVPHVTEKYGQNWTSPENMVTSGPKLKERIPNEKYVFEKNNKYYD
SNEVELEEITFYTTNDSSTAYKMYENEELDAIFGSIPPDLIKNLKLRSDYYSSAVNAIYFYAFNTHIKPLDNVKIR
KALTЛАIDRETLTYKVLNGTPTRRATPNFSSYAKSLELFNPEIAKTLAEAGYPNGNGFPILKLKYNTNEAN

TABLE 1. Nucleotide and Amino Acid Sequences

KKICEFIQNQWKKNLNIDVELENEEWTTYLNTKANGNYEiarAGWIGDYADPLTFLSIFTQGYTQFSSHNSNPEY
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t606.aa

CCNNKERKEGVSKISLGAEPSSLDPQLAEDNVASKMIDTMFRGIVTGDPTGGNKPGLAKGWDISSDGTVYTFNL
REKITWSDGVAITAEGIRSKYRLIRLNKETGSKYVEMVSKVIKNGQKYFDGQVTDSELGIRAIDEKTLEITLESPKP
YFIDMLVHQSFIPVPHVTEKYGQNWTSPENMTSGPKLKERIPNEKYVFEKNNKYYDSNEVELEEIFYTTNDS
STAYKMYENEELDAIFGSIPPDLIKNLKLRSDYYSSAVNAIYFYAFNTHIKPLDNVKIRKALTIAIDRETLTYKVL
DNGTTPTRRATPNFSSSYAKSLELFNPEIAKTLAEAGYPNGNGFPIKLKYNTNEANKKICEFIQNQWKKNLNID
DVELENEEWTTYLNTKANGNYEiarAGWIGDYADPLTFLSIFTQGYTQFSSHNSNPEYNELIKKSDLELDPIKRQ
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f606.nt

ATGAAATTACAAAGGTCAATTATTTTAATAATATTTCTAACTTTCTTGTATAATAACAAGGAAAGAAAAG
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GCAAAAGGGTGGGATATTCTCTGATGGAACAGTTACACATTAACTTAACCTAACAGAGAAAAAAATCACTGGAGTGACG
GAGTTGCAATCACTGCAAGGAATTAGAAAATCTTATCTAGAACATTAAAGAAAATGGCTCAAAGTACGT
TGAAATGGTTAAATCGGTAAATTAAAATGGTCAAAATTTGATGGACAAGTGACTGACTCTGAACATTGGAATT
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GAGCAACTCCCAACTTATGTTCAATTCTATGCAAAAGTTAGAATTATTAACCTGAAATTGCAAAACCCCT
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AAAAAAATTGTAATTATTCAAAACCAATGGAAAAAAATTAAATTGATGTGGAACTTGAAAACGAAGAAT
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TCCTTGACATTAAAGCATATTCAACACAAGGATACACACAATTCTCATCTCATATTACTCAAACCCAGAATAC
AACGAACATTATAAAGAAATCCGACCTTGAGCTTCAATAAAAGACAAGACATTAAAGACAAGCAGAAGAGA
TAATTATTGAAAAAGATTTCCTAACAGCACAATACATATATGGAACAGTTACCTTTCAAGAAATGACAATG
GACAGGGTGGAACACCAATATTAGAAAGATTGATTCTCAGCTAAATTAAAAATAAATAA

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TGTTGTAATAACAAGGAAAGAAAAGAAGGAGTATCATTAAAATAAGCTGGGAGCAGAGCCAAGCAGTCTGACC
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GATGTGGAACCTGAAAACGAAGAATGGACAACATACTTAAACACTAAGGCAAATGAAATTGAAATAGCAAGAG
CAGGATGGATAGCGGATTATGCTGATCCTTGACATTAAAGCATATTCAACACAAGGATACACACAATTCTCATC

TABLE 1. Nucleotide and Amino Acid Sequences

TCATAATTACTCAAACCCAGAATACAACGAACCTATAAAGAAATCCGACCTTGAGCTTGATCCAATAAAAAGACAA
GACATTTAAGACAAGCAGAAGAGATAATTATTGAAAAAGATTTCATAGCACCAATATACATATATGGAAACA
GTACCTTTCAAGAAATGACAAATGGACAGGGTGGACACCCAATATTAGAAAGATTGATTCTCAGCTAAA
ATAAAAAAATAAATAA

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MFNRSSCVLQNFLFLFLSLVSCFAKKEISGNNFIAHSKEFDLNNLNWLWNFDYTKKNFDKHFNIDPSSYIYVA
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KKNDKALSLLNKLDKMKFSDYQENENILLKAVLYLNLSNVSESKIYFNELFENLPANYLHVRAYDYFIIENKSRYF
GANFLNLVRFKYEVANGNFNGAINILNKNGLNDYYDNNIVLSDVYKAFISSGKVSNALTFFSKIJKSKYKNYYLGIL
NLREKNNLGLLLKEYLEGLDLNNEINRLDNLNTAFSNLIFTKSARDYFAESLPKFYTEGDKKNSTFIKILEEYIL
ESIQLEDYGNLYKLYSNAQKVISNSVLSKAFINARLIYHKLIKPNVSGEYKSLLHSAVNYDKWSYSSFMSRYLLD
QNIDEFFTGGSDIKYEQSDYEIFLEGFLKFNLNCNYVRGFISEDFRNGYKFSLDFYRKVYDELLKSENYYDATLVIN
YLVNQDESALMENDYKRLYPYLYGSLIEYWAKRRGLEASVVFSLIKAESSFEKNAVSKPGAVGLMQVMPSTANDIS
KELKYFNYDLKIPKDNIIIGTYYLKKRISTTGSPLYKALASYNGGIGNVRKWEKSYGHLSEKELFIEAIPFSQTRNYI
KKILVYSVFYDALYEKKGIDSIVKIMGEFPKNZ

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CFAKKEISGNNFIAHSKEFDLNNLNWLWNFDYTKKNFDKHFNIDPSSYIYVAYLFKKIGFEKFVEYMKKAIANG
DSIASQFAGIKLIEYFNSAKEYFASELIGEKLYKKYENNKFIIILGYFKSLYQWQKKNDKALSLLNKLDKMKFSDYQE
NENILLKAVLYLNLSNVSESKIYFNELFENLPANYLHVRAYDYFIIENKSRYFGANFLNLVRFKYEVANGNFNGAI
NILNKNGLNDYYDNNIVLSDVYKAFISSGKVSNALTFFSKIJKSKYKNYYLGILNLREKNNLGLLLKEYLEGLDLN
NEINRLDNLNTAFSNLIFTKSARDYFAESLPKFYTEGDKKNSTFIKILEEYILESIQLEDYGNLYKLYSNAQKVIS
NSVLSKAFINARLIYHKLIKPNVSGEYKSLLHSAVNYDKWSYSSFMSRYLLDQNIDEFITGGSDIKYEQSDYEIF
LEGFLKFNLNCNYVRGFISEDFRNGYKFSLDFYRKVYDELLKSENYYDATLVINYLVNQDESALMENDYKRLYPYLY
GSLIEYWAKRRGLEASVVFSLIKAESSFEKNAVSKPGAVGLMQVMPSTANDISKELKYFNYDLKIPKDNIIIGTYY
LKKRISTTGSPLYKALASYNGGIGNVRKWEKSYGHLSEKELFIEAIPFSQTRNYIKKILVYSVFYDALYEKKGIDSVI
VKIMGEFPKNZ

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ATGTTAATAGAAGTTCTTGTATTACAAAATTTCTTTCTTTTATTTTAAGTTAGTTCTTGCTTTG
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CAAGAGTCTTGTCTGCTTAATTGATAAAATGGCTTATTCTCATTGAGTGTAGTACTTATTAGAT
CAAATATTGATGAATTGTTACAGGTGGGCTGATATTAGTGTAGGCAATCCGATTATGAGATTGTTGAG
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TATCTTGTAAATCAAGATGAATCTGCTTTAATGGAGAATGACTATAAAAGACTTATCCTTATTGATGATCTT
TGATAGAATATTGGCTAAAGGAGAGGGCTGAGCTAGTGTGATTGTTAATAAGCAGAGAGTAGCTT

TABLE 1. Nucleotide and Amino Acid Sequences

TGAAAAAAATGCTGCTCAAAACCGGGTGCCTGGCCTATGCAGGTTATGCCATCAACAGCAAATGATATTC
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 AAAAAAAATATTAGTTATTCGGTATTTATGATGCTTGATGAAAAGAAGGAATAGATTCAAGTAATAGTTAAAA
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TGCTTGCAAAAAAGAAATCTCAGGCAATAATTTATTAAGGCGCATTCAAAAGAGTTGATTTAAATAATTAA
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 GATAGCATTGCATCCCAGTTGCTGGGATTAAGCTTATTGAATATTAACTCAGCAAAGAGATTGGCATCTG
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 GGATCTTGATAGAATATTGGCTAAAGGAGAGGGCTGAGCTAGTGTGATTCTTAAATAAAGCAGAGA
 GTAGCTTGAAAAAAATGCTGCTCAAAACCGGGTGCCTGGCCTTATGCAGGTTATGCCATCAACAGCAAATGA
 TATTCTAAAGAACTTAAGTATTTAACTATGATTAAAGATCAGGAAAGATAATATAATAATGGAACATATTAT
 TTAAAAAAAGAATATCTACAACGGCAGTCTTATAAGGCTCTTGCCTTATAATGGGGTATTGGTAATGTTA
 GAAAGTGGAGAAAAGTTATGGACATTTGCAAAAGAGCTTTTATTGAGGCAATTCCCTTAGTCAAACAGGA
 TTATATAAAAAAATATTAGTTATTCGGTATTTATGATGCTTGATGAAAAGAAGGAATAGATTCAATA
 GTTAAAATTATGGCGAATTCCCCAAAATTAA

f11-12.nt

TAAAAGGAGA ATATTTTAT GAGAAAAAGT TTGTTTTAT ATGCATTATT AATGGGAGGA
 TTGATGTCTT GTAATCTAGA TTCCAAATTAA TCTAGTAACA AAGAACAAA AAATAACAAT
 AATGTAAG AAGTTTCGGA TAGTGTCAA GAAGATGGTC TTAATGATT ATATAATAAT
 CAAGAAAAGC AAAAAGCTT TACTAAAAT TTTGGAGAAC GGAAATATGA GGATTAAATT
 AATCCTATAAG AGCCTATAAT ACCTTCAGAA TCACCAAAGA ATAAGGCTAA TATACCAAAAT
 ATTTCATTG CGCATACTGA AAAAAGAG AAAAAAAGG AGAATTAAAT CCCTTCTACT
 AATGAAGAAA AGGAAGCTGA TGCAAGCAATT AAATATTAG AAGAAAATAT TCTTAAAAC
 TCTAAATTCTGAAATTAAAT TAGAGAAGTA CGTGTAAATTAA AGATGAATA TGCTTTAATA
 AAAGCTGATT TGTATGATGT ATTGGAAAG ATTAACAATA AAAAACATC ATTAATGGAG
 AATCCTAAGA ACAATAGAGA TAAGATAAAAT AAATTAACAC AATTGTTGCA AAATAATTAA
 AAGATAGATA GTGAACCTGA GCAGCTTATA AATATGATTG ATATGGCAGA AAATGAAATA
 AGCTCTGGGG CTTCTTTTGACAAACGCT CAGAAAAGGT TAAAAGAAAG CATTATTAA
 AGATTAGAGA GTAAAAATAA TAGATCTTAT GCATTAATG TGTCTAGACA GGCTTTAAGT
 GACGCAAGGA GTGCTTTAAG TAATTTAGAA TCTTTGCCT CTAAAAGAAT TGAACCAATG
 GTGAGAAAGG AAGAAATAA AGAGCTTATT AAACATGCAA AAACTGTTT AGAAAGTCTC
 AATAAAAAT AA

TABLE 1. Nucleotide and Amino Acid Sequences

t11-12.nt

TTGTAATCTAGATTCCAAATTATCTAGTAACAAAGAACAAAAATAACAATAATGAAAAGAAGTTCGGATAGT
 GTTCAAGAAGATGGCTTAATGATTATAATAATCAAGAAAAGCAAAAAGCTTACTAAAAATTGGAGAAC
 GGAAATATGAGGATTTAATTAACTCTATAGGCCTATAATACCTTCAGAACATCACCAGAATAAGGCTAATATACC
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 ATTAAAATTGTCTAGACAGGCTTAAAGTGACGCAAGAAGTGCTTAAAGTAATTAGAATCTTTGCCTCTAAAAGA
 ATTGAACCAATGGTGAGAAAGGAAGAAATAAGAGCTTATTAAACATGCAAAACTGTTTAGAAAGTCTCAATA
 AAAAA

f11-12.aa

KENIFMRKSL FLYALLMGGL MSCNLD SKLS SNKEQKNNNN VKEVSDSVQE DGLNDLYNNQ
 EKQKSFTKMF GERKYEDLIN PIEPIIPSES PKNKANIPNI SIAHTEKKET KKENLIPSTN
 EKEADAAIK YLEENILKNS KFSELIREVR VIKDEYALIK ADLYDVIGKI NNKKTSLMEN
 PKNNRDKINK LTQLLQNNLK IDSELEQLIN MIDMAENEIS SAAFFFDNAQ KRLKESIICKR
 LESKNNRSYA LKLSRQALSD ARSALS NLES FASKRIEPMV RKEEIKELIK HAKTVLES LN
 KK

t11-12.aa

CNLD SKLSSNKEQKNNNNVKEVSDSVQEDGLNDLYNNQE KQKSFTKMF GERKYEDLIN PIEPIIPSES PKNKANIP
 NISIAHTEKKET KKENLIPSTN EKEADAAIK YLEENILKNS KFSELIREVR VIKDEYALIK ADLYDVIGKINNKK
 TSLMENPKNNRDKINK LTQLLQNNLK IDSELEQLIN MIDMAENEIS SAAFFFDNAQ KRLKESIICKR
 LESKNNRSYA LKLSRQALSD ARSALS NLES FASKRIEPMV RKEEIKELIK HAKTVLES LN
 KK

f11-4.nt

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 ATCACAAAAT TAACTCCGGA AGAGCTAGAA AATTAGCAA AGGAAGCTCA AGATGACTCT
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 GTAAAGGATA CTCCTCGCTT AATCAAATTG ATAAAGAATT CATCAGAAAA AATTGATTG
 GTTTTCAAA CACTAATTAA TATAGGTAT AATGCTACCT ATGCAGCCAA AAGTAATTG
 AAGAATGGAC TAAAGATGGT GAAATTACTG GATGAGTTGC TAAAATATC GTAAAGTAGC
 AATGGTGATA AAAGTACCCCA AAAATACAAT GAACTTAAA CCGTTCTAAA TAAGTTTAAT
 GCTGAAAATT CGGTAAGCGT TTCTTTAAA GAACATTCAA ACAGTAAAAT TGAAACTAAA
 AAATGTATTG AAACCTTTAT GAAAAATGTA GAAACATACT TTGAAGGTGT ATGCAGCGAA
 CTTAAAACA AAAATGATGG TGAGTACGAA AAAACATTGA CAACTTAAAG CTAA

t11-4.nt

ATGTAAATGGTATGTAGACAATACCATTGATGAAGCAACTGTAGAAAGTAAATCAGCACTAACATCTATTGATCAA
 GTATTAGATGAGATAAGTGAAGGCCACAGGCCCTAAGTCGGAAAAAAATCACAAAATTAACTCCGGAAGAGCTAGAAA
 ATTTAGCAAAGGAAGCTCAAGATGACTCTGAAAAATTCCAAAAAGAAATTGAGATCAAAAATACCAAGGAAAG
 TAAAACATAGAAGTAAAGGATACTCCTCGCTTAACTCAAATTGATAAGAATTGATCAGAAAAAATTGATTCGGTT
 TTTCAAACACTAATTAAATAGTTATAATGCTACCTATGCAAGCCAAAAGTAATTGAGAATGGACTAACAGTGG
 TGAAATTACTGGATGAGTTGCTAAAATATCGGTAAGTAGCAATGGTATAAAAGTACCCAAAATACAATGAAC
 TAAAACCGTTGTAATAAGTTAATGCTGAAAATTGGTAAGCGTTCTTTAAAGAACATTCAAACAGTAAAATT

TABLE 1. Nucleotide and Amino Acid Sequences

GAAACTAAAAATGTATTCAAACCTTATGAAAATGTAGAAACATACTTTGAAGGTGTATGCAGCGAACTTAAAAA
ACAAAAATGATGGTGACTACGAAAAA

f11-4.aa

RSLQMSKLIL AISILLIISC KWYVDNTIDE ATVESKSALT SIDQVLDEIS EATGLSSEKI
TKLTPEELEN LAKEAQDDSE KSKKEIEDQK NTKESKNIEV KDTPRLIKLI KNSSEKIDS
FQTLINIGYN ATYAAKSNLK NGLKMKVLLD ELLKISVSSN GDKSTQKYNE LKTVVNKFNA
ENSVSVSFKE HSNSKIETKK CIQTLMKNVE TYFEGVCSEL KNKNDGEYEK TLTTLS

t11-4.aa

CKWYVDNTIDEATVESKSALTSIDQVLDEISEATGLSSEKITKLTPEELENLAKEAQDDSEKSKEIEDQKNTKES
KNIEVKDTPRLIKLIKNSSEKIDSVFQTLINIGYNATYAAKSNLKNGLKMVKLLDELLKISVSSNGDKSTQKYNE
KTVVNKFNAENSVSVSFKEHSNSKIETKKCIQTLMKNVETYFEGVCSELKNKNDGEYEK

f112-1.nt

TGAATCTCTA AAGATTTAG CAGGGGAGAA AATATGAAAA AAAGTTTTT ATCAATATA
ATGTTAATT CAATAAGTTT ATTATCATGT GATGTTAGTA GATTAATCA GAGAAATATT
AATGAGCTTA AAATTTTGT TGAAAAGGCC AAGTATTATT CTATAAAATT AGACGCTATT
TATAACGAAT GTACAGGGAGC ATATAATGAT ATTATGACTT ATTCCGAAGG TACATTTCT
GATCAAAGTA AGGTTAATCA AGCTATATCT ATATTTAAAAA AAGACAATAA AATTGTTAAT
AAGTTTAAGG AGCTTGAAAA GATTATAGAA GAATACAAAC CTATGTTTT AAGTAAATT
ATTGATGATT TTGCGGGATC CGTT

t112-1.nt

ATGTGATGTTAGATTAAATCAGAGAAATTAAATGAGCTAAAATTTTGTGAAAAGGCCAAGTATTATTCT
ATAAAATTAGACGCTATTATAACGAATGTACAGGGAGCATATAATGATATTGACTTATCGGAAGGTACATT
CTGATCAAAGTAAGGTTAATCAAGCTATATCTATATTAAAAAAGACAATAAAATTGTTAATAAGTTAAGGAGCT
TGAAAAGATTATAGAAGAATACAAACCTATGTTTAAGTAAATTGATGATT

f112-1.aa

ISKDFSRGEN MKKSFLSIYM LISISLLSCD VSRLNQRNIN ELKIFVEKAK YYSIKLDIA
NECTGAYNDI MTYSEGTFSQ QSKVNQAISI FKDNKIVNK FKELEKIIIE YKPMFLSKLI
DDFAGSV

t112-1.aa

CDVSRLNQRNINELKIFVEKAKYYSIKLDIAYN
ECTGAYNDIMTYSEGTFSQSKVNQAISIFKKDNKIVNKFKEL
EKIIIEYKPMFLSKLIDDF

f14-8.nt

TAAATACAGA GCCATTCAAG GAGAGTATTT ATGAAATACT ATATATGTGT GTGTGTTTT
TTGCTTTGAA ATGCTTGAA TTCAGATTT AGCACTAATC AAGAAGATAT TAAATATCCA
TCTGATAAAG AGAAATCAA ATCCAACATG GAAGCAAGCT CTAAGAAGA AGATCCAAAT
AAAAAAATAA AAAATACACT GCTTAATGAT TTAATAAATT TGATAGAAAT AGCTAATGAG
CATAAAGAAA AATATGAAA AAGAATGCAA GAAGAACCTT CAGATCAATA CGGAATATTG
GCTTTCCAGG AATTAGACTT GTCCGTTGGA AAAATATCTG AAGACACCCC GCAATCTAA
AAATTAGAA AAAACACCTA TTCTCCCTTA AGCGCTATTG ATGTCAATAA ATAAAAGAT
CTTTCAGAGA TTATAAGAAA TTCCGGCCAA ATACAAGGTT TATTTAATAT TTTCAACAGA
TTCCGGAGGCA TTTTGACGA CTCACCTAAT CACGTATATT CTAAAAAGA TATCCTAGGG
GGACTAGAAA TTTGGATTT AGATAAACTA AAAAATTCTG TTGAAAATT ACTATCTATA

TABLE 1. Nucleotide and Amino Acid Sequences

AAAGAAACTT TCTCAAAAAT GCTAAATCAA CTTTTATTAG ATTATAAAAA TGATAAAAGAT
 CATATACGAA CAGAGACAAA TAAACTTAAA TCTCATACAA CTGCACTTT CGAACAACTT
 GATAAAAAG AAGACGAAGC ATATGAACCT AAAAATCAGA TATTTCAAT AAGTAACCTT
 TAA

t14-8.nt

TTGCAATTAGATTTAGCACTAATCAAGAAGATATTAAATATCCATCTGATAAAAGAGAAATCAAATCAAACATG
 GAAGCAAGCTCTAAAGAAGAAGATCAAATAAAAAAATACACTGCTTAATGATTTAATAAATTGATAG
 AAATAGCTAATGAGCATAAGAAAAATATGAAAAAGAATGCAAGAAGAACCTTCAGATCAATACGGAATATTGGC
 TTTCCAGGAATTAGACTTGTCCGTTGGAAAAATATCTGAAGACACCCCGCAATCTAAAAAATTAGAAAAAACACC
 TATTCTCCCTTAAGCGCTATTGATGTCAATAAATTAAAGATCTTCAGAGATTATAAGAAATCGGGCAAATAC
 AAGGTTTATTTAATATTTCAACAGATTGGAGGCATTTGACGACTCACTTAATCACGTATATTCTAAAAAGA
 TATCCTAGGGGACTAGAAATTGGATTAGATAAACTAAAAAATTGTTGAAAAATTACTATCTATAAAAGAA
 ACTTTCTCAAAATGCTAAATCAACTTTATTAGATTATAAAAGATCATATACGAACAGAGACAAATA
 AACTTAAATCTCATACAACACTGCACTTTCGAACACTTGATAAAAAGAAGACGAAGCATATGAACCTAAAAATCA
 G

f14-8.aa

IQSHSRRVFM KYYICVCVFL LLNACNSDFS TNQEDIKYP DKEKSNSNME ASSKEEDPNK
 KIKNTLLNDL INLIEIANEH KEKYEKRME EPSDQYGILA FQELDLSVGK ISEDTPQSKK
 FRKNTYSPLS AIDVNKLKDL SEIIRNSGQI QGLFNIFNRF GGIFFDSLH VYSKKDILGG
 LEILDLDKLK NSFEKLLSIK ETFSKMLNQL LLDYKNDKDH IRTETNKLKS HTTALFEQLD
 KKEDEAYEPK NQIFSISNL

t14-8.aa

CNSDFSTNQEDIKYP DKEKSNSNME ASSKEEDPNKKIKNTLLNDL INLIEIANEH KEKYEKRME EPSDQYGILA
 FQELDLSVGK ISEDTPQSKK FRKNTYSPLS AIDVNKLKDL SEIIRNSGQI QGLFNIFNRF GGIFFDSLH VYSKKD
 ILGGLEILDLDKLKNSFEKLLSIK ETFSKMLNQL LLDYKNDKDH IRTETNKLKS HTTALFEQLD
 KKEDEAYEPK NQIFSISNL

f17-6.nt

TAAAGGAGGG TATTATGAA ATACCACATA ATTACAACTA TATTTGTTT TCTGTTTTA
 GCTTGCAGGC CGGATTTAA TATCGATCAA AAAGACATTA AATACCCGCC TACTGAAAAA
 TCAAGGCCA AAACGTAAAG CTCTAAGCAA AAAGAATCAA AGCCTAAAAC AGAAGAAGAG
 CTTAAGAAAA AACACAACA AGAAGAGCTT AAGAAAAAC AACACAACAAG AAGAAGAGAA AGAGCTTAAG
 AAAAACAAAC AAGAAGAAGA GCTTAAGAAA AAACAACAAG AAGAAGAGAA GGAAGAACTA
 AGAAAACAAC AACTAAAAAA TACGCTATCT AATGATTAA AAAAGCAAAT AGAATCGGCC
 TACAATTTA AAGAAAAATA TGTAAAAGT ATGGAAAAAG AACCTGAAGA CCATTACGGG
 ATGACGTCTT TTAGGGGATT GAATTGGGG CCAGGGACTG AAGATATATC TGACAATACC
 GAAAGATCTA TAAGATATAG AAGACACACT TATACTGTT TAAGCCCCCT GGATCCTCAT
 GAATTAAAGG AATTGCAAA TATTATCAA GATATAAATA AACTAGCATC AGTAGCAAGT
 ATATTTAATT CTTTAGCGC TATTGGAGGA GCTCTTGACA TAGTAAGTGA TCACCTATAT
 TTCAAAAAAG ACAATCTAGA CAAACTAGAT ATTCAGATT TAGAAATACT TAAAAATTCA
 TTTGAACAAA TATTATATAT AAAAGGAAGT GTTGCAGGAA AAGCAAAAAA ACTTTTATTA
 GATTATAAAA ATCTAAAAAC AGATATTAAT AAGCTTAAAT CTTATCAAA TGAACGGTT
 AATGGAATTAA AGCAACAAGC TCTAGAAGCA GAAAATCTAG AAGAGCTTAT AGTGTCAAAA
 TATAAACTTT AA

t17-6.nt

TTGCAGGCCGGATTTAATATCGATCAAAAGACATTAATACCCGCCACTGAAAAATCAAGGCCAAAATGAA
 AGCTCTAAGCAAAAGAATCAAAGCCTAAACAGAAGAAGAGCTTAAGAAAAACACAAGAAGAGCTTAAGA

TABLE 1. Nucleotide and Amino Acid Sequences

AAAAACACAAGAAGAAGAGCTTAAGAAAAAAACAACAAGAAGAAGAGCTTAAGAAAAAAACAACAAGAAGAGAGA
GGAAGAAGACTAAGAAAACAACAACTAAAAAATACGCTATCTAATGATTAAAAAAGCAAATAGAATCGGCCTACAAT
TTTAAAGAAAAATATGTAAAAGTATGGAAAAGAACCTGAAGACCATTACGGGATGACGTCTTTAGGGGATTGA
ATTGGGGCCAGGGACTGAAGATATATCTGACAATACCGAAAGATCTATAAGATATAGAAGACACACTTATACTGT
TTTAAGCCCCCTGGATCCTCATGAATTAAAGGAATCGCAAATATTCAAGATATAAAACTAGCATCAGTA
GCAAGTATATTAAATTCTTTAGCGCTATTGGAGGAGCTTGACATAGTAAGTGTACCTATATTCAAAAAAG
ACAATCTAGACAAACTAGATATTGCAAGATTAGAAACTTAAATTCATTGAAACAAATATTATATAAAAGG
AAAGTGTGCAAGGAAAGCAAAAAACTTTATTAGATTAAAAATCTAAAACAGATATTAAAGCTTAAATCT
TATTCAAATGAACTGGTTAATGGAATTAGCAACAAGCTCTAGAAGCAGAAAATCTAGAAGAGCTTATAGTGTCAA
AAATATAAAACTT

f17-6.aa

RRVFMKYHII TTIFVFLFLA CRPDFNIDQK DIKYPPTEKS RPKTESSKQK ESKPKTEEEL
KKKQQEEELK KQQQEEELKK KQQEEELKKK QQEEEKEELR KQQLKNTLSN DLKKQIESAY
NFKEKYVKSM EKEPEDHYGM TSFRGLNWGP GTEDISDNT E RSIRYRRHTY TVLSPLDPHE
LKEFANIIQD INKLASVASI FNSFSAIGGA LDIVSDHLYF KKDNLKDLDI ADLEILKNSF
EQILYIKGSV AGKAKKL禄D YKNLKTDINK LKSYSNELVN GIKQQALEAE NLEELIVSKY
KL

t17-6.aa

f19-2.nt

TAAAGAAAGA TAAATCATA TTCAAGGAGA GTATTTATGA AACACTATAT AATTGTGCAT
ATATTGTTT TTCTATTTT AAATGCTTGT TATCCAGTTG CATCTAATAA AATAGAAATTA
AACCTAAAA CAGAAACAAG CTTAAATCAA GAAGAAGTCC CAAATCAAGA AGCAAACATAC
AAAGAAGAAA AAGAAGCAAA AGAAGAAGGC ATTAATAAAA AACAGAAAA CACGCTGCTT
AATGATTTAA GAAATTAAAG AGAAACAGCT AAAAAAGATA ATGATAAATA TACACAAAAG
TTAAAAGAAG AATCCTCAAG CC_AATACGGA ATACTGGCTT TCAAAGATT GTTCTGGCTA
GATGGAACAA ATGAACAATT GTCCCGCAAAT ACCGAAAGAT CTAAGCCTA TAGAAAACGA
GCTTATAGCA TCTTAAATAC TATTAATGAC GCTTCCTTAA AGAATTTC AGAAATTGTA
ATGGCATCAG GACAAACACA GGGCATATTT AATACCCCTA ACTCACTTGG GGGTAATTAA
GAAAAGATAG TTAAATTGTTT GTATCCAAA AAAGACAATT TGGAAAATT AGAGACTTCA
GTTTAAAAAA AGCTTAAAGA TTCTTGAA AATTTTTAG AGATAAAAAA AATCGCCTCA
GAAATGATGC ACAAGCTTT ATTAGACTAT CAAAATAATA CAAATCGTAT ACAAAACAGAT
AAAAATGAAC TTAAGTCTTA TGCAGACACA CTTTTCAATC AAATGACAAA AAAACCCGAA
GAAGCACTAA AGCTAAAAAA TACCATATGC TCAATAGAGG ACCTTTAA

t19-2.nt

TGTTATCCAGTGCATCTAATAAAATAGAATTAAAACCTAAACAGAAACAAGCTTAAATCAAGAAGAAGTCCC
AATCAAGAAGCAAACCTACAAAGAAGAAAAAGAAGCAAAAGAAGAAGGCTTAATAAAAAAACAGAAAACAGCTGC
TTAATGATTTAAGAAATTAAATAGAAACAGCTAAAAAGATAATGATAAAATACACAAAAGTTAAAAGAAGAATC
CTCAAGCCAATACGGAATCTGGCTTCAAAGATTGTTCTGGCTAGATGGAACAAATGAAACAATTGTCGCAAAT
ACCGAAAGATCTAAAGCCTATAGAAAACGAGCTTATAGCATCTTAAATACTATTAAATGACGCTTCTTAAAGAATT
TTTCAGAAATTGTAATGGCATCAGGACAAACACAGGGCATATTAAACCCCTTAACTCACTTGGGGTAATTGAA
AAAGATAGTTAATTGTTGTATCCCAAAAAGACAATTGAAAAATTAGAGACTCAGTTTAAAGCTTAA
GATTCTTGGAAAATTAGAGATAAAAAAAATCGCCTCAGAAATGATGCACAAAGCTTATTAGACTATCAA
ATAATACAAATCGTATACAAACAGATAAAAATGAACCTAAGTCTTATGCAGACACACTTTCAATCAAATGACAAA
AAAACCCGAAGAAGCACTAAAG

TABLE 1. Nucleotide and Amino Acid Sequences

f19-2.aa

RKIKSYSRRV FMKHYIIVHI FVFLFLNACY PVASNKIELK PKTETSLNQE EVPNQEANYK
 EEEKEAKEEGI NKKTENTLLN DLRNLIETAK KDNDKYTQKL KEESSSQYGI LAFKDLFWLD
 GTNEQLSANT ERSKAYRKRA YSILNTINDA SLKNFSEIVM ASGQTQGIFN TLNSLGGNFE
 KIVNCLYPKK DNLEKLETSV LKKLKDSLLEN FLEIKKIASE MMHKLLLQYQ NNTNRIQTDK
 NELKSYADTL FNQMTKKPEE ALKLKNTICS IEDL

t19-2.aa

CYPVASNKIELKPKTETSLNQEEVPNQEANYKEEKEAKEEGINKKTENTLLNDLRNLIETAKKDNDKYTQKLKEES
 SSQYGILAFKDLFWLDGTNEQLSANTERSKAYRKRAYSLNTINDASLKNFSEIVMASGQTQGIFNTLNSLGGNFE
 KIVNCLYPKKDNLEKLETSVLKKLKDSLLEN FLEIKKIASEMMHKLLLQYQ NNTNRIQTDKNELKSYADTLFNQMTK
 KPEEALK

f19-4.nt

TAATCTATAC TAATTGAGGA GAATATTTT ATGAAAAACA ACATAATTTT ATGCATGTT
 GTTTTTTAC TTTTAAATAG CTGCACCGCT AACCATGAAG CTGAAGCGAA ATAAAAAAA
 CATGTTGATA AAACAAAAAA CGAATATATT AATGAAATAA AAAATTTAAT AGCAACAAACC
 AAAGAAATCA TCGAAAAACG AAAATTGCTA CAAGCTAAC CAGTAGATCA AAACCCCGTA
 GATGATACAA ACAATAAGAA AGTTTCGAG ATAGATAAAA GAGCTTCGA TTTTATAAAT
 AGTTTTTAA CAGATGATGA ATTTAATAAA TTGTAACAA TATTCATCAA ACCAACACTA
 AAATCACCCG GAAAAGTATT AAATAGCATA GCAATTCTAG AGCTAACAT AGAGCAGGTA
 ATTAATCACC TAGACTAAA AAATGAGACC TTAATAAAG CAAGCTCTT AGATTGGAA
 AAGATCAAA ATCCCTTGA ACAGCTGTC TCTATAAGGA ATTTTTTTC ACAATCATA
 AAAAGGGTCT TATTAGATCA TCAAAACAAT GAAAATTCTA TAAAACCAGA TGATTCTAAA
 TCAGGAACCT ATTTGATAC GATATAGGAT CAGTTAATG AAAAAATAA AGAGGTTAGA
 AATCTGAAAA AAACCATATT ATCACTGCCG AATTAA

t19-4.nt

CTGCACCGCTAACCATGAAGCTGAAGCGAAAATAAAAAACATGTTGATAAAACAAAAACGAATATATTAAATGAA
 ATAAAAAATTTAATAGCAACAACCAAAAGAAATCATGAAAAACGAAAATTGCTACAAGCTAAACCAGTAGATCAAA
 ACCCGTAGATGATAAAACAATAAGAAAGTTTCGAGATAGATAAAAGAGCTTCGATTTATAAATAGTTTTT
 AACAGATGATGAATTAAATTTGTAACAATATTCTATAAAACCAACACTAAATCACCGGGAAAAGTATTAAAT
 AGCATAGCAATTCTAGAGCTAACATAGAGCAGGTAAATTACACCTAGACTCAAAAAATGAGACCTTAAATAAG
 CAAGCTCTTAGATTGGAAAAGATCAAAATCCCTGAACAGCTGTTCTCTATAAGGAATTTTTCACAAAT
 CATAAAAGGGTCTTATTAGATCATCAAAACAATGAAAATTCTATAAAACCAGATGATTCTAAATCAGGAACCTAT
 TTCGATACGATATACGATCAGTTAATGAAAAAATAAGAGGTTAGAAATCTGAAAAAA.

f19-4.aa

SILIEENIFM KNNIILCMCV FLLLNSCTAN HEAEAKIKKH VDKTKNEYIN EIKNLIATTK
 EIIIEKRKLLQ AKPVDQNPVD DTNNKKVFEI DKRAFDINS FLTDDEFNKF VTIFHKPTLK
 SPGKVLNSIA ILELNIEQVI NHLD SKNETL NKASSLDLEK IKN SLEQLFS IRNFFSTI
 RVLLDHQNNE NSIKPDDSKS GTYFDTIYDQ FNEKNKEVRN LKKTILSLPN

t19-4.aa

CTANHEAEAKIKKHVDKTNEYINEIKNLIATTKEIIEKRKLLQAKPVDQNPVDDTNKKVFEIDKRAFDINSFL
 TDDEFNKFVTIFHKPTLKSPGKVLNSIAILELNIEQVINHLD SKNETLNKASSLDLEK IKN SLEQLFS IRNFFSTI
 IKRVLLDHQNNENS IKPDDSKSGTYFDTIYDQFNEKNKEVRN LKK

f19-6.nt

TABLE 1. Nucleotide and Amino Acid Sequences

TAAAGGAGAG TATTAATGAA ATGCCATATA ATTGCAACTA TATTTGTTTT TCTATTTTA
 GCTTGCAGTA CAGATTTAA TACTGATCAA AAAGGCATTA AATACCCGCC TACCGAAAAA
 TCAAAGCCC AAACTGAAGA CTCTAAGCAA AAAGAATTAA AGCCTAAAAC AGAAAAAAGAA
 CTAAGAAAAA AACAACAAC TAAAAATAAA CTACTTAATG ATTTAAAAAA TTCAATAGAA
 ACAGCTAATA AGCATAAAAGA AAAGTATAAA AAAAGAATGA AAGAAGAACC CGAAGATCAA
 TACGGGGTAC AGGCTTCAGA AGGATCGAAT TGGGGGCCGG GGACTGAAGA TGTATCTGCC
 AACACCGAAA GATCTATAAG ATTTAGAAGA CATACTTATA CTATTTAAG CACGCTGAGT
 CTCATGAAT TAAAGGAATT CTCAAATATT GTTACAAATG AAAATAAACT GGTGCCAGTA
 GTAGATATGT TTAATTTCTT TAGCTCTATT GGGACAGCTC TTGATATAAC AACCGATAGC
 TTATATCCC AAAAGACAAT CTGGACAAAC CAGATCTGTC GGATTTAG

t19-6.nt

TTGCAGTACAGATTTAATACTGATCAAAAGGCATTAATACCCGCCTACCGAAAAATCAAAGCCC AAAACTGAA
 GACTCTAACGCAAAAGAATTAAAGCTAAAACAGAAAAAGAACTAAAGAAAAACAACACTAAAAAATAAACACTAC
 TTAATGATTAAAAAATTCAATAGAACAGCTAATAAGCATAAAAGAAAAGTATAAAAAAAGAATGAAAGAAGAAC
 CGAAGATCAATACGGGGTACAGGCTTCAAGGATCGAATTGGGGCCGGGACTGAAGATGTATCTGCCAACACC
 GAAAGATCTATAAGATTAGAACATACCTATACTATTAAAGCACGCTGAGTCTTCATGAATTAAAGGAATTCT
 CAAATATTGTTACAAATGAAAATAAACTGGTGCCAGTAGTAGATATGTTAATTCTTAGCTCTATTGGACAGC
 TCTTGATATAACAAACCGATAGCTTATATCCAAAAAGACAATCTGGACAAACAGATCTGTCGG

f19-6.aa

RRVLMKCHII ATIFVFLFLA CSTDFNTDQK GIKYPPTEKS KPKTEDSKQK ELKPKEKEL
 KKKQQLKNKL LNDLKNSIET ANKHKEKYKK RMKEEPEDQY GVQAFKGSNW GPGTEDVSAN
 TERSIRFRRH TYTILSTLSL HELKEFSNIV TNENKLVPVV DMFNFFSSIG TALDITTDLS
 YPKKTIWTNQ ICRI

t19-6.aa

CSTDFNTDQKG IKYPPTEKS KPKTEDSKQKELKPKTEKELKKQQLKNKLNDLKNSIETANKHEKYKKRMKEEP
 EDQYGVQAFKGSNW GPGTEDVSANTERSIRFRRHTYTLSTLSL HELKEFSNIVTNEKLVPPVDMFNFFSSIGTA
 LDITTDLSLYPKKTIWTNQICR

f21-4.nt

TAGGAGACAA TCTTATGAA TAAAAAATA AAAATGTTA TTATTTGTGC TATTTTTATG
 CTGATAAGTT CTTGTAAGAA TGATGTAAC AGTAAAGATT TAGAAGGGC GGTGAAAGAT
 TTAGAAAGTT CAGAACAAA TGAAAAAAA ACAGAACAAAG AGATAAAAAA ACAAGTTGAA
 GGATTTTAG AAATTTAGA GACAAAAGAT TTAAACACAT TAGATACAAA AGAAATTGAA
 AAACAAATTG AAGAATTAAA GAATAAGATA GAAAATTAG ACTCTAAAAA AACTTCTATT
 GAAACATATT CTGGGTATGA AGAAAAATA AACAAAATA AAGAAAATT AAACGGAAA
 GGACTTGAAG ATAAATTAAA TGAACTTCA GAGAGCTTAA AAAAGAAAAA AGAGGAGAGA
 AAAAAAGCTT TACAAGAGGC TAAAAAGAAA TTGAGAGAT ATAAAAACCA AGCTGAATCT
 GCAACTGGAG TAACGCATGG TTCTCAAGTC CAAAGACAAAG GTGGTGTGG ATTACAAGCT
 TGGCAGTGTG CTAATAGTTT GGGGTTAAA AATATGACTA GTGGTAATAA TACTAGCGAT
 ATGACCAATG AAGTTATAAC TAATCGCTT AAAAGATTG AAGAAGAACT TAAAAATATT
 GGAGAAACTG TAAGAGTAA AAAAGAATAA

t21-4.nt

TTGTAAGAATGATGTAACTAGTAAAGATTAGAAGGGCCGGTGAAGAGATTAGAAAGTTCAAGAACAAAATGTAAAAA
 AAAACAGAACAAAGAGATAAAAAAAACAAGTGAAGGATTTAGAAATTAGAGACAAAAGATTAAACACATTAG
 ATACAAAAGAAATTGAAAAACAAATTCAAGAATTAAACAATAAGATAGAAAATTAGACTCTAAAAAAACTTCTAT
 TGAAACATATTCTGGGTATGAAGAAAAATAACAAAATTAACGGAAAAGGACTTGAAGATAAA

TABLE 1. Nucleotide and Amino Acid Sequences

TTAAATGAACCTTCAGAGAGCTTAAAAAGAAAAAGAGGAGAGAAAAAGCTTACAAGAGGCTAAAAGAAAT
 TTGAAGAGTATAAAACCAAGCTGAATCTGCAACTGGAGTAACCCATGGTCTCAAGTCCAAAGACAAGGTGGTGT
 TGGATTACAAGCTGGCAGTGTGCTAATAGTTGGGTTAAAAATATGACTAGTGGTAATAATACTAGCGATATG
 ACCAATGAAGTTATAACTAATTGCTTAAAAGATTGAAGAAGAACTTAAAATATTGGAGAACTGTAGAAGGTA
 AAAAGAA

f21-4.aa

ETIFMNKKIK MFIICAIFML ISSCKNDVTS KDLEGAVKDL ESSEQNVKKT EQEIKKQVEG
 FLEILETKDL NTLDTKEIEK QIQELKNKIE KLDSSKTSIE TYSGYEEKIN KIKEKLNGKG
 LEDKLNESEL SLKKKKEERK KALQEAKKKF EYKNQAES A TGVTGHSQVQ RQGGVGLQAW
 QCANSLGFKN MTSGNNTSDM TNEVITNSLK KIEEELKNIG ETVEGKKE

t21-4.aa

CKNDVTSKDLEGAVKDLESSEQNVKKTEQEIKKQVEGFLEILETKDLNTLDTKEIEKQIQLKNKIEKLDSSKTSI
 ETYSGYEEKINKIKEKLNGKGLEDKLNESELSSLKKKKEERKKALQEAKKKFEYKNQAESATGVTHGSQVQRQGGV
 GLQAWQCANSLGFKNMTSGNNTSDMTNEVITNSLKKIEEELKNIGETVEGKKE

f24-1.nt

TAAGCTGGTA ACACGTAAA GACAGCTGAG GGGGCTTCAA GTGGTACTGA TGCAATTGGA
 GAAGTTGTGG ATAATGATGC TAAGGTTGCT GATAAGGCGA GTGTGACGGG GATTGCTAAG
 GGGATAAAGG AGATTGTTGA AGCTGCTAGG GGGAGTGAAG AGCTGAAAGT TGCTGCTGCT
 AAAGAGGGCA ATGAAAAGGC AGGGAAGTTG TTTGGGAAGG CTGGTGCCTA TGCTCATGGG
 GACAGTGAGG CTGCTAGCAA GGCAGCTGGT GCTGTTAGTG CTGTTAGTGG GGAGCAGATA
 TTAAGTGCCTA TTGTTAAGGC TGCGGATGCG GCTGAGCAGG ATGGAAGAA GCCTGCAGAT
 GCTACAAATC CGATTGCTGC TGCTATTGGG AATAAAAGATG AGGATGCGGA TTTTGGTGT
 GGGATGAAGA AGGATGATCA GATTGCTGCT GCTATTGCTT TGAGGGGGAT GGCTAAGGAT
 GGAAAGTTG CTGTGAAGAA TGATGAGAAA GGGAAAGGCTG AGGGGGCTAT TAAGGGAGCT
 GCTGCAATTG GAGAAAGTTGT GGATAATGCT GGTGCTGCGA AGGCTGCTGA TAAGGATAGT
 GTGAAGGGGA TTGCTAAGGG GATAAAAGGAG ATTGTTGAAG CTGCTGGGG GAGTGAAAG
 CTGAAAGCTG CTGCTGCTGA AGGGGAGAAT AATAAAAGG CAGGGAAGTT GTTTGGGAAA
 GTTGATGGTG CTGCTGGGA CAGTGAGGCT GCTAGCAAGG CGGCTGGTCC TGTTAGTGC
 GTTAGTGGGG AGCAGATATT AAGTGCATT GTTAAGGCTG CTGGTGAGGC TGAGCAGGAT
 GGAGAGAACG CTGAGGATGC TAAAAATCCG ATTGCTGCTG CTATTGGAA GGGTAATGGG
 GATGGTGCCTT AGTTGATCA GGATGAGATG AAGAAGGATG ATCAGATTGC TGCTGCTATT
 GCTTGAGGG GGATGGCTAA GGATGGAAAG TTTGCTGTGA AGGGTAATAA TGAGAAAGAG
 AAGGCTGAGG GGGCTATTAA AGAAGTTAGC GAGTTGTTGG ATAAGCTGGT AACAGCTGTA
 AAGACAGCTG AGGGGGCTTC AAGTGGTACT GATGCAATTG GAGAAGTTGT GGATAATGNT
 GCNAAGGNTG CTGATAAGGC GAGTGTGACG GGGATTGCTA AGGGATAAAA GGAGATTGTT
 GAAGCTGCTN GGGGGAGTGA AAAGCTGAA GTTGCTGCTG CTANAGNGN NAATAATAAA
 GAGGCAGGGAG TTGTTGG GAAGGCTGGT GCTGATGCTA ATGGGGACAG TGAGGCTGCT
 AGCAAGGCGG CTGGTGCCTG TAGTGTGTT AGTGGGGAGC AGATATTAAAG TGCGATTGTT
 AAGGCTGCGG CTGCTGGTGC GGCTGATCAG GATGGAGAGA AGCCTGGGA TGCTAAAAT
 CCGATTGCTG CTGCTATTGG GAAGGGTAAT GCGGATGATG GTGCGGATTT TGGTGATGGG
 ATGAAGAAGG ATGATCAGAT TGCTGCTGCT ATTGCTTTGA GGGGGATGGC TAAGGATGGA
 AAGTTGCTG TGAGAAAGGA TGAGAAAGGG AAGGCTGAGG GGGCTATTAA GGGAGCTAGC
 GAGTTGTTGG ATAAGCTGGT AAAAGCTGTA AAGACAGCTG AGGGGGCTTC AAGTGGTACT
 GCTGCAATTG GAGAAAGTTGT GGATAATGCT GCGAAGGCTG CTGATAAGGA TAGTGTGACG
 GGGATTGCTA AGGGGATAAA GGAGATTGTT GAAGCTGAG GGGGGAGTGA AAAGCTGAAA
 GTTGCTGCTG CTAAAGGGGA GAATAATAA GGGCAGGGAG TTGTTGG GAAGGCTGGT
 GCTAATGCTC ATGGGGACAG TGAGGCTGCT AGCAAGGCGG CTGGTGCCTGT TAGTGTGCTGTT
 AGTGGGGAAAC AGATATTAAAG TGCGATTGTT AAGGCTGCTG GTGAGGCTGC TGGTGATCAG
 GAGGGAAAGA AGCCTGAGGA GGCTAAAAT CCGATTGCTG CTGCTATTGG GGATAAAAGAT
 GGGGATGCGG AGTTAATCA GGATGGGATG AAGAAGGATG ATCAGATTGC TGCTGCTATT

TABLE 1. Nucleotide and Amino Acid Sequences

GCTTTGAGGG GGATGGCTAA GGATGGAAAG TTTGCTGTGA AGGATGGTGG TGAGAAAGAG
 AAGGCTGAGG GGGCTATTAA AGGAGTTAGC GAGTTGTTGG ATAAGCTGGT AAAAGCTGTA
 AAGACAGCTG AGGGGGCTTC AAGTGGTACT GCTGCAATTG GAGAAGTTGT GGCTGATGCT
 GCTAAGGTTG CTGATAAGGC GAGTGTGACG GGGATTGCTA AGGGGATAAA GGAGATTGTT
 GAAGCTGCTG GGGACAGTGA GGCTGCTAGC AAGGCAGCTG GTGCTGTTAG TGCTGTTAGT
 GGGGAGCAGA TATTAAGTGC GATTGTTAAG GCTGCGGCTG CTGGTGCAGGC TGAGCAGGAT
 GGAGAGAAGC CTGCAGAGGC TAAAAATCCG ATTGCTGCTG CTATTGGAA GGGTGATGGG
 GATGCGGATT TTGGTGAGGA TGGGATGAAG AAGGATGATC AGATTGCTGC TGCTATTGCT
 TTGAGGGGGA TGGCTAAGGA TGGAAAGTTT GCTGTGAAGA ATGATGAGAA AGGGAAGGCT
 GAGGGGGCTA TTAAGGGAGC TGCTGCAATT GGAGAAGTTG TGGATAATGC TGGTGCTGCG
 AAGGCTGCTG ATAAGGATAG TGTGAAGGGG ATTGCTAAGG GGATAAAAGGA GATTGTTGAA
 GCTGCTGGGG GGAGTGAAGAA GCTGAAAGCT GCTGCTGCTG AAGGGGAGAA TAATAAAAAG
 GCAGGGAAAGT TGTTTGGAA AGTTGATGGT GCTGCTGGGG ACAGTCAGGC TGCTAGCAAG
 GCGGCTGGTG CTGTTAGTGC TGTTAGTGGG GAGCAGATAT TAAGTGCAGTG TTAAAGGCT
 GCGGATGCGG CTGAGCAGGA TGGAAAGAAG CCTGCAGATG CTACAAATCC GATTGCTGCT
 GCTATTGGGA ATAAAGATGA GGATGCGGAT TTTGGTGTG GGATGAAGAA GGATGATCAG
 ATTGCTGCTG CTATTGCTTT GAGGGGGATG GCTAAGGATG GAAAGTTG TGTAAGGGT
 AATAATGAGA AAGGAAGGC TGACGGGGCT TCAAGTGGTA CTGATGCAAT TGGAGAAGTT
 GTGGATAATG ATGCGAAGGC TGCTGATAAG GCGAGTGTGA CGGGGATTG TGAGGGGATA
 AAGGAGATTG TTGAAGCTGC TGAGGGGGAGT GAAAAGCTGA AAGCTGTTGC TGCTGCTACA
 AGGGAGAATA ATAAAGAGGC AGGGAAAGTTG TTGGGAAAG TTGATGATGC TCATGCTGGG
 GACAGTGAGG CTGCTAGCAA GGCGGCTGGT GCTGTTAGTG CTGTTAGTGG GGAGCAGATA
 TTAAGTGCCTA TTGTTACGGC TGCGGCTGCT GGTGAGCAGG ATGGAGAGAA GCCTGCAGAG
 GCTACAAATC CGATTGCTGC TGCTATTGGG AAGGGTAATG AGGATGGTGC GGATTTGTT
 AAGGATGAGA TGAAGAAGGA TGATCAGATT GCTGCTGCTA TTGCTTGAG GGGGATGGCT
 AAGGATGGAA AGTTGCTGT GAAGAGTAAT GATGGTGAGA AAGGGAAGGC TGAGGGGGCT
 ATTAAGGAAG TTAGCGAGTT GTTGGATAAG CTGGTAAAGAC AGCTGAGGGG
 GCTTCAAGCG GTACTGATGC AATTGGAGAA GTTGTGGCTA ATGCTGGTGC TGCGAAGGGCT
 GCTGATAAGG CGAGTGTGAC GGGGATTGCT AAGGGGATAA AGGAGATTGT TGAAGCTGCT
 GGGGGAGTA AAAAGCTGAA AGCTGCTGCT GCTGAAGGGG AGAATAATAA AAAGGCAGGG
 AAGTTGTTG GGAAGGCTGG TGCTGGTGC GGTGCTAATG GGGACAGTGA GGCTGCTAGC
 AAGGCGGCTG GTGCTGTTAG TGCTGGTTAG

t24-1.nt

TGGTGAAGGCTGAGCAGGATGGAGAGAAGCCTGAGGATGCTAAAAATCCGATTGCTGCTGCTATTGGGAAGGGTAAT
 GGGGATGGTGCAGGAGTTGATCAGGATGAGATGAAGAAGGATGATCAGATTGCTGCTGCTATTGCTTTGAGGGGGAA
 TGGCTAAGGATGGAAAGTTGCTGTGAAGGGTAATAATGAGAAAGAGAAGGCTGAGGGGGCTATTAAAGAAGTTAG
 CGAGTTGTTGGATAAGCTGGTAACAGCTGTAAGACAGCTGAGGGGGCTCAAGTGGTACTGATGCAATTGGAGAA
 GTTGTGGATAATGNTGCNAAGGGNTGCTGATAAGGCCAGTGTGACGGGGATTGCTAAGGGGATAAAGGAGATTGTT
 AAAGCTGCTGGGGAGTGAAGGCTGAAAGTTGCTGCTGCTANAGNGNNAAATAATAAGAGGCAGGGAAAGTTGTT
 TGGGAAGGCTGGTGTGCTAATGGGACAGTGAGGCTGCTAGCAAG

f24-1.aa

AGNTVKTAEG ASSGTDIAIGE VVDNDAKVAD KASVTGIAKG IKEIVEAARG SEKLKVAAK
 EGNEKAGKLF GKAGANAHGD SEAASKAAGA VSAVSQEQL SAIVKADAA EQDGKKPADA
 TNPIAAAIIGN KDEDADFGDG MKKDDQIAAA IALRGMAKDG KFAVKNDEKG KAEGAIKGAA
 AIGEVVDNAG AAKAADKDSV KGIAGKIKEI VEAAGGSEKL KAAAEGENN KKAGKLFKGKV
 DGAAGDSEAA SKAAGAVSAV SGEQILSAIV KAAGEAEQDG EKPEDAKNPI AAAIIGKGN
 GAEFDQDEMKG KDDQIAAAIA LRGMAKDGK AVKGNNEKEK AEGAIKEVSE LLDKLVTAVK
 TAEGASSGTD AIGEVVDNAXA KXADKASVTG IAKGIKEIVE 'AAXGSEKLKV AAAXXXNNKE
 AGKLFKGAGA DANGDSEAAS KAAGAVSAVS GEQILSAIVK AAAAGAADQD GEKPGDAKNP
 IAAAIGKGNA DDGADFGDGM KKDDQIAAAI ALRGMAKDGK FAVKKDEKGK AEGAIKGASE
 LLDKLVKA
 TAEGASSGTA AIGEVVDNAA KAADKDSVTG IAKGIKEIVE AAGGSEKLKV
 AAKGENNKG AGKLFKGAGA NAHGDSEAAS KAAGAVSAVS GEQILSAIVK AAGEAAGDQE

TABLE 1. Nucleotide and Amino Acid Sequences

GKKPEEAKNP IAAAIGDKDG DAEFNQDG MK KDDQIAAAIA LRGMAKDGF AVKDGG EKEK
 AEGA IKGVSE LLDKLV KAVK TAEGASSGTA AIGEVVADAA KVADKASVTG IAKGIKEIVE
 AAGDSEAASK AAGAVSAVSG EQILSAIVKA AAAGAAE QDG EKPAEAKNPI AAAIGKGDGD
 ADFGEDGMK K DDQIAAAIAL RGM AKDGFKA VKNDEKGKAE GAIK GAAAIG EVVDNAGAAK
 AADKDSVKG I AKGIKEIVEA AGGSEKLKAA AAEGENNKA GKLFGKVDGA AGDSEAASKA
 AGAVSAVSGE QILSAIVKA DAAE QDGK P ADATNPIAAA IGNKDEDADF GDGMKKDDQI
 AAAIALRGMA KDGKFAVKG N EKGKAEGAS SG TD AIGEVV DNDAAKADKA SVTGIAKG IK
 EIVEAAGGSE KLKAVAAATR ENNKEAGKLF GKVDDAHAGD SEAASKAAGA VSAVSGEQIL
 SAIVTAAAAG EQDGEKPAEA TNPIAAAIGK GNEDGADFGK DEMKKDDQIA AAAIALRGMAK
 DGKFAVKSND GEKGKAEGAI KEVSELLDKL VKAVKTAEGA SSGTDAIGEV VANAGAAKAA
 DKASVTGIAK GIKEIVEAAG GS KKLKAAA EGENNKKAGK LFGKAGAGAG ANGDSEAASK
 AGAVSAG

t24-1.aa

GEAEQDGK PEDAKNPIAAAIGKNGDGAEFDQDEM KKDDQIAAAIALRGMAKDGF AVKG NNEKEKAEGAIKEVS
 ELLDKLVTAVKTAEGASSGTD AIGEVVVDNXAKXADKASVTG IAKGIKEIVEAAXGSEKLK VAAAXXXNNKEAGKLF
 GKAGADANGDSEAASK

f28-2.nt

TAAAAAGGAA ATATAAAATAT TATGCGATT A TGTTAATAA AAATTTTAT TATACTTAAT
 TTAGTATT A GTTCTCTTT TTTATTTGAA AGTTGTTCTG GTTTCTATC TAAAAAATCT
 ATAGAACAGT TTGCATTAGC ATTAAAAGAT CATCAAGAAA ATAAAATACT TACTAATACT
 TCACTAGATA AAAATAGTAA GGAAATTGAA TCTCCTAAAG ACGTTACATC ATCAAATAAA
 AAAACTTATG ATCCAATCTT ACAAGTAGGT TCTAATCAC AC ATATGTCAGA TGATCCTGGT
 GCTAATAATA AAGAATCCCT ACCAAATTCA AGTCCAGCAA TAATACAAAA TGACTCGCAT
 GCTAAAATA ATGTAAGGAT GGAAGAAAAT AAATCAGCTA CTCCACACAA TGATCCAATT
 GAACAAAGTA ATTTAAAAA TAGCCTTAAC ACAACAAGTA AAACCTCTGC TATTCTTCA
 GAAGAAGAAA TTAAAGCTAA CTTAGATGAA TTTGCACAAG AAGAGTATGA GCAAACATCT
 CTTTCAGAAA TTAAAATGC CACGCAAATT GTTAATCATG CTAATCCTGA AAACAAATT
 AACAAATACAC TCCTTGAGTT TGAAAAGAT TATGAAAATT TATCAAAC TTATTCTCT
 AATTTAGACG CATCTCCTT GAATAGAAAA ATAAAGACTA TTATGCCTAA ATTACAAGAA
 ATGCGTTCTT TTATGGAGCA AGCAACTAAT TCTGGGTAT CTGCTAAAGG CATGCTAGAT
 GAGGCTAAGG ATAAAATAGC AGAATCTATT TATAAAAGAC TATACATGG CAATTCTAC
 CGGTTCGGTG GCAGTTTAA CGGACGTGAT ATGCAACATG CAAAAAATT AGCATAACAGA
 GCTATAGACT TTGCTTCTGC ATGCATTGAA TATACACAAA AAGCTATTGA TTATCTTCA
 CAGGGAAATT CTTGCAAAAAA AGAAATAGAA AATATATTCA AGCTTTAA

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AAAAGATCATCAAGAAAATAAAACTACTAATAC TCTCAGTAGATAAAAATAGTAAGGAAATTGAATCTCTAAA
 GACGTTACATCATCAAATAAAAACCTTATGATCCAATCTTACAAGTAGGTTCTAATCAACATATGTCAGATGATC
 CTGGTCTAATAATAAAAAGAATCCCTACCAAATTCAAGTCCAGCAATAATACAAATGACTCGCATGCTAAAATAA
 TGTAAGATGGAAGAAAATAATCAGCTACTCCACAACATGATCCAATTGAACAAAGTAATT TAAAGCTTAAAGCCTT
 ACTACAACAAGTAAAACCTCTGCTATTCCCTCAGAAGAAGAAAATTAAGCTAACTTAGATGAATTGCAACAGAAG
 AGTATGAGCAAACATCTCTTCAAGAAATTAAAATGCCACGCAAATTGTTAATCATGCTAATCTGAAAACAAATT
 AAACAATACACTCCTTGAGTTGAAAAGATTATGAAACTTTATCAAAC TTGTATTCTCTAATTAGACGCATCT
 CCTTTGAATAGAAAATAAGACTATTATGCTAAATTACAAGAAAATGCGTTCTTTATGGAGCAAGCAACTAATT
 CTTGGGTATCTGCTAAAGGCATGCTAGATGAGGCTAAGGATAAACTAGCAGAATCTATTATAAAAAGACTATACAA
 TGGCAATTCAACCGGTTCGGTGGCAGTTAACGGACGTGATATGCAACATGCAACAAATTAGCATAACAGAGCT
 ATAGACTTTGCTTCTGCATGCAATTGAATATACACAAAAGCTATTGATTATCTTCAACAGGGAAATTCTTGCAAA
 AAGAAAATAGAAAATATATTCAAG

f28-2.aa

TABLE 1. Nucleotide and Amino Acid Sequences

KGNINIMRLC LIKIFIIPNL VFSSLFLFES CSGFLSKKSI EQFALALKDH QENKNTTNTS
 VDKNSKEIES PKDVTSSNKK TYDPILQVGS NQHMSDDPGA NNKESLPNSS PAIIQNDSHA
 QNNVKMEENK SATPQHDPIE QSNFKNSLTT TSKTPAIPSE EEIKANLDEF AQEEYEQTSL
 SEIKNATQIV NHANPENKLN NTLLEFEKDY ETLSNLLFSN LDASPLNRKI KTIMPKLQEM
 RSFMEQATNS WVSAGMLDE AKDKLAEsiY KRLYNGNSYR FGGSFNGRDM QHAKNLLAYRA
 IDFASACIEY TQKAIDYLQQ GNSCKKEIEN IFKL

t28-2.aa

KDHQENKNTTNTSVDKNSKEIESPKDVTSSNKKTYDPIQVGSNQHMSDDPGANNKESLPNSSPAIIQNDSHAQNN
 VKMEENKSATPQHDPIEQSNFKNSLTTSKTPAIPSEEEIKANLDEFAQEEYEQTSLSEIKNATQIVNHANPENKL
 NNTLLEFEKDYETLSNLLFSNLDASPLNRKIKTIMPKLQEMRSFMEQATNSWVSAGMLDEAKDKLAEsiYKRLYN
 GNSYRFGGSFNGRDMQHAKNLLAYRAIDFASACIEY TQKAIDYLQQGNSCKKEIENIFK

f28-3.nt

TAGATGAATT TAATTGCTAA ATTATTTATT TTATCCACTT TAGTTCAAT TCCAAATATC
 CTCTCTTGTAA ACCTATATGA TAATCTTGCAGAACACGCTG AGCAGGTTAC AGACATACTA
 GACAACAAACA AGTCTTTAA TACTTTAGGA AGCAGCAATG AGAGTAGAAG TCGCAGGCT
 AGAAGTACAA ATAATGCTTA TATGAAACAA AACATAGACAA AAAATCATTT AGTTGTTGCA
 GATATGCAAA ATGATAATAG TAGCAGCAGT CTTCCCCAAC AAGTTAATAG TGAATCCAGT
 AAAGCTAATG AAGATAGTAA TATTATGAAG GAAATTGAAT CTTCTACAGA AGAGTGCCT
 AGACTAAGAA AAGATTTAGA AACTATAAAA CAAATACTTG ATAATATAGA AAGCTTGCTT
 AATAACAGCTA ATTCTTATTT AGAGAACGCT AGAAAAGCAC CTAAATCTAA TCAAGATAAT
 CAAACCTTAT TGCTTAGCCT GCACCAAGCT ATTGCTAAGG TTAAGAGTAG TCATACTTCT
 TTTATCATTGTTATAATGA TGCAATTAAAT TCCCTGGAA TAGCTGATAC TGCCTTTAAA
 GATGCAAAGA GAAAGGCACT TGAGGCATAA

t28-3.nt

TTGTAACCTATATGATAATCTTGCAGACAACGCTGAGCAGGTTACAGACATACTAGACAAACAAGTCTTTAAT
 ACTTTAGGAAGCAGCAATGAGAGTAGAAGTCGCAGGCCTAGAAGTACAAATAATGCTTATATGAAACAAAACATAG,
 ACAAAAATCATTTAGTTGTCAGATATGCAAAATGATAATAGTAGCAGCAGTCTTCCCCAACAAAGTTAATAGTGA
 ATCCAGTAAAGCTAATGAAGATAGTAATATTATGAAGGAAATTGAATCTCTACAGAAGAGTGCCTAGACTAAGA
 AAAGATTTAGAAACTATAAAACAAACTTGATAATATAGAAAGCTTGCTTAATACAGCTAATTCTTATTTAGAGA
 ACGCTAGAAAAGCACCTAAATCTAATCAAGATAATCAAACCTTATTGCTTAGCCTGCACCAAGCTATTGCTAAGGT
 TAAGAGTAGTCATACTTCTTTATCATTGTTATAATGATGCATTAAATTCCCTGGAAATAGCTGATACTGCCTT
 AAAGATGCAAAGAGAAAGGCAGTTGAGGCA

f28-3.aa

MNLIAKLFIL STLVSIPNIL SCNLYDNLAD NAEQVTDILD NNKSFTNLGS SNESRSRRPR
 STNNAYMKQN IDKNHLVVAD MQNDNSSL PQQVNSESSK ANEDSNIMKE IESSTEECAR
 LRKDLETIKQ ILDNIESLLN TANSYLENAR KAPKSQDNQ TLLLSLHQAI AKVKSSHTSF
 IICYNDAFNS LGIADTAFKD AKRKAVEA

t28-3.aa

CNLYDNLADNAEQTIDLDNNKSFTLGSSNESRSRRPRSTNNAYMKQNIDKNHLVVADMQNDNSSLQQVNSE
 SSSKANEDSNIMKEIESSTEECARLRKDLETIKQILDNIESLLNTANSYLENARKAPKSQDNQTLSSLHQAIAKV
 KSSHTSFIICYNDAFNSLGIADTAFKDARKAVEA

f31-2.nt

TAAAAAAATA AGGAGGTATT AATGAAAAGG AAAAGCAATA TATGTATTTC ACTTCTAGTC
 ACAATATTAT TTGTGTCTTG CAAGTTTTT GGAAATAAAA GCGCAAGTAA AGAAAAAGAA

TABLE 1. Nucleotide and Amino Acid Sequences

GAAACTTCTT TTTCTGATAC TGCTAGCAAG ATTAGTAAGT CGGGAACAGC TGCTTCTTCA
 GACAAACAAAG AAAAAAAATAC AAGTGATGTT ACAGGTGACG CCAAAAAGCA TACTAGTAGC
 CCTTACATGC TTGCTGATGC CCTTATTGTT AGTGATACTA CTAATAGAGA TAGAGATAAG
 CAAGAAAATA AAGATAAAATT AAATGAAGAA GATAAAAAAA AGCTTAATGC TTTTTTTAGC
 ACAACTAAAA CATATCAATC TAGCCTAGAT TCCATTATA ACAAAATATAC AGGCTATTAT
 AATACCATTG ATACCTATGG CAGCTGTGAT ACGTATCGCA TTGAGTGTGTT TAGTGTAGGA
 CCTTCTGAAA AACGTAACAA AGCTCTTGCT GATCTAGAGA AGTTAAAAGT AGACGAAAAG
 TACACTCAGC TTAGCACAAT GTTAAAGAGT GCTGTGCCA GTTATTACAA AAAAAATTAA
 GATGATTCTA TTGCACAGTA TAAGGAAGCC ATAAAGCAGG CTATTGAAGC TGAAAGTAAA
 ATAGAGACAG TAAAAGACTA TGCAACAGCT CAAAGTGTG CCGATGACGA AAAGAAAAGA
 AATATAGATA ATTTAAAAAT AGTTAGAGAT GTTCTTCTTA TTATTAAAAA AACTATTGAG
 AAAGCCAGCC GATCTTATGC TGATGCTTT GCTATTGCAA CATCTAGCTT ATCTTGAGC
 GAATTTAAGC AAGCTGTTAA AGAGTTAAT GATGCTGCTA AACAAATATGC TAATGGAAAT
 AAAGGAGACA ATGCTGTCAA TGTATTGTA GGCACATTCTT CTAGTATGCC TTATGTCAAA
 TTAAAGATG AGTTGCAAG AGCAAAAATG TTTGCTCGTA ATTATAGAGG AGACGAGGT
 GACAAGATGA TAAGAGCTAT CGACAGCTG TGTGATGTTT ATAAAAAAAGT TCGCCTTTAG

t31-2.nt

TTGCAAGTTTTGGAAATAAAAGCGCAAGTAAAGAAAAAGAAGAAACTTCTTTCTGATACTGCTAGCAAGATT
 AGTAAGTCGGAACAGCTGCTCTTCAGACAAACAAAGAAAAAAATACAAGTGATGTTACAGGTGACGCCAAAAGC
 ATACTAGTAGCCCTTACATGCTGCTGATGCCCTATTGTTAGTGATACTACTAATAGAGATAGAGATAAGCAAGA
 AAATAAAAGATAAAATTAAATGAAGAAGATAAAAAAAAGCTTAATGCTTTTTAGCACAACATAAAACATATCAATCT
 AGCCTAGATCCATTATAACAAATATACAGGCTATTATAATACCATTGATACCTATGGCAGCTGTGATACGTATC
 GCATTGAGTGTGTTAGTGAGGACCTCTGAAAAACGTAACAAAGCTTGTGATCTAGAGAAGTTAAAAGT
 CGAAAAGTACACTCAGCTTAGCACAATGTTAAAGAGTGCTGTGCCCTAGTTATTACAAAAAAATTAGATGATTCT
 ATTGCACAGTATAAGGAAGCCATAAGCAGGCTATTGAGCTGAAAGTAAAATAGAGACAGTAAAAGACTATGCAA
 CAGCTCAAAGTGTGCCGATGACGAAAAGAAAATAGATAATTAAAAATAGTTAGAGATGTTCTCTTAT
 TATTAAAAAAACTATTGAGAAAGCCAGCCGATCTATGCTGATGCTTTGCTATTGCAACATCTAGCTTATCTTGT
 AGCGAATTAAAGCAAGCTGTTAAAGAGTTAATGATGCTGCTAAACAAATATGCTAATGAAATAAGGAGACAATG
 CTGTCATGTTATTGAGGACTATTCTAGTATGCCTTATGTCAAATTAAAGATGAGTTGCAAGAGCAAAAT
 GTTGCTCGTAATTATAGAGGAGACGAGGTAGACAAGATGATAAGAGCTATCGACAAG

f31-2.aa

KNKEVLMKRK SNICISLLVT ILFVSKFFG NKSASKEKEE TSFSDTASKI SKSGTAASSD
 KQEKNTSVDT GDAKKHTSSP YMLADALIVS DTTNRDRDKQ ENKDKLNEED KKKNNAFFST
 TKTYQSSLDS IYNKYTGYYN TIDTYGSCDT YRIECFSVGP SEKRKQALAD LEKLKLDEKY
 TQLSTMLKSA VPSYYKKNLD DSIAQYKEAI KQAI EAESKI ETVKDYATAQ SAADDEKRN
 IDNLKIVRDV LLIKKTIEK ASRSYADAF A IATSSLSCSE FKQAVKEFND AAKQYANGNK
 GDNAVNVIVG TISSMPYVKF KDEFARAKMF ARNYRGDEV KMIRAIKLC DVYKKVAL

t31-2.aa

CKFFGNKSASKEKEETSFSDTASKISKSGTAASSDKQEKNTSDVTGDAKKHTSSPYMLADALIVSDTTNRDRDKQE
 NDKKLNEEDKKKLNAAFFSTTKTYQSSLDSIYNKYTGYYNTIDTYGSCDTYRIECFSVGPSEKRKQALADLEKLKLD
 EKYTQLSTMLKSAVPSYYKKNLD DSIAQYKEAI KQAI EAESKI ETVKDYATAQ SAADDEKRNIDNLKIVRDV
 LLIKKTIEKASRSYADAF A IATSSLSCSE FKQAVKEFND AAKQYANGNK
 GDNAVNVIVG TISSMPYVKF KDEFARAKMF ARNYRGDEV KMIRAIKLC DVYKKVAL

f32-4.nt

TAAGGAAATA TGAGGAATAT TAGCAATTGT ATCAAATATA TTATATTAAC AATGCTTATT
 GGATTATTAA TTTTTGTTG TGCAACCTTT GTTGGTTGA TTGGAATTAA TTATTCAAAT
 AACTTTAAAG AAGAGCGGAA TTATTCAATA AGCCCAATAG ATAGTGTAT TATGCCGTAAA
 TGTTATTAA AAGAATTAA GTCTGGACTT ATTAAAAGCG TATTCTTTAA GAAATTAGAT

TABLE 1. Nucleotide and Amino Acid Sequences

GTAATGTTA ACTCTAAAAA TTTAAGGAG CAAATAAGG TAGATAAACAAATCTGCTA
 AATTCTTATC CATCTTATCA TATGGAGTTT GTCGTAGTT ATAATGGATT TTTAATGAAT
 TTAAAAATG TTATTTTAA TGGTATAGAT GATGCTAAAT TATACGATCA ACGTGATATG
 GTTACGGAG GATTTAGATA CTCAAAAGAG GCTTATTTCC AAATTATTGG CAATTATGAT
 GTAAATTAA ATAAAATGAA ACAATATACT CCAGCAATTG TAGTAAATGT TTTCAAAATT
 AACATTAATG ATGCTTTATT TAACTCGTTA TTAAAGCAAA AAACCTTAAAGTTACTTTG
 ATTTCCCATATAAAGA GTATATTAA CAAACTAATA ATTTCTTATC AAAGTATAAT
 TTCAACAC CAGAAAAGGA GAATAGTCT TACTAA

t32-4.nt

AAATAACTTTAAAGAAGAGCGGAATTATTCAATAAGCCAAAGATAGATGTATTATGCGTAAATGTTATTTAAA
 GAATTTAAGTCTGGACTTATTAAAAGCGTATTCTTAAAGAAATTAGATGTAATGTTAACTCTAAATTTAAGG
 AGCTAAATAAGGTAGATAACAAAATCTGCTAAATTCTTATCCATCTATCATGGAGTTGCGTAGTGATAA
 TGGATTTAAATGAATTAAAATGTTATTTAAATGGTATAGATGATGCTAAATTATACGATCAACGTGATATG
 GTTACGGAGGATTTAGATACTCAAAGAGGCTTATTCCTAAATTATGGCAATTATGATGTTAAATTAAATAAAA
 TGAACAAATATACTCCAGCAATTGAGTAAATGTTCAAAATTAAACATTAATGATGCTTATTTAACTCGTTATT
 AAAGCAAAACTTTAAAGTTACTTGATTTCCCATATAAATAAAAGAGTATATTTACAAACTAATAATTCTTA
 TCAAAGTATAATTTCACACCCAGAAAAGGAGAATAGTTCTTAC

f32-4.aa

GNMRNISNCI KYIILMLIG LLIFCCATFV WLIGIFYSNN FKEERNYSIS PIDSVIMRKCFYKEFKSGLI
 KSVFFKKLDV NVNSKNFKEL NKVDKQNLLN SYPSYHMEFV VVDNGFLMNF
 KNVIFNGIDD AKLYDQRDMV YGGFRYSKEA YFQIIGNYDV KLNKMKQYTP AIVVNVFKIN
 INDALFNSLL KQKTLKVTLI SHNKEYILQ TNNFLSKYNF QTPEKENSSY

t32-4.aa

NNFKEERNYSISISPDSVIMRKCFYKEFKSGLI KSVFFKKLDVNNSKNFKELNKVDKQNLLNSYPSYHMEFV VVDN
 GFLMNFKNVIFNGIDDAKLYDQRDMVYGGFRYSKEAYFQIIGNYDV KLNKMKQYTP AIVVNVFKIN INDALFNSLL
 KQKTLKVTLISHNNKEYILQ TNNFLSKYNF QTPEKENSSY

f4-15.nt

TAAATGAGCA AAAAGTAAT TTTAATATTA CTAGAAATT TGATCTTGTCTTGTGATTAA
 TCTATAATAA AAGAACAAA AACCAAAGAA AAAACATCTG AAAAGCAAGA ATCTGAAAAAA
 CAAAATATTG AAAACAAAGA GCCTGAAAAA CAGAAACAAA ATGCAGCAAA ATAATCCCT
 ACGGTATCAA TTCAACGGT AGAAATAAGG GAATCAAATC AAATTCCAAA AAGCATTGAG
 AAGTACTACA AGCAAGCTT TCCGATTCAA ACATTCACTC TTGATTTAG CATCACAAGA
 GAAAAGGAAT TTCTAAACC AGAAGATAAA ATCTTGCCCA CACAGGGAA AGTGGAGCT
 TTGAGCATCT TAATAAATAA AAAATTGTTA GACTTTAAAG CCCCAGAAAA TCCAAAAAGC
 TCAACTTTAA AAAATTCAA AGAAATTAAA AATATTGAGA ATTTCTTCCA AAATCAAGAC
 TTATTATTG TCTTAAACCCT TAAAGATAAA AATAACAACA AACTATTAA CATCATGCTC
 AATCCCCAA ACGACATCCA AAAACCCAAA GATTATATT TAAAAGACCT TAAAGACACA
 ATTAAAAAGG GTACTGGTGA GAAATACTTA AATCCTATCT ATAGATTCA AATAAAAAAC
 AAAAGATT ATCATTCAAT AGATTACAAC AAAGTGACTA TTAGCGAAAA AACAATAGAA
 TTGGACCTAC TGCCTCACGA ACAAGTCTT CAAATGAATA AAAATTTCAC TAAAATTAA
 GACACAATAA CAGACTTAA TAATCTAAAAA TTAGTAATTC AAAAGAATT AGTGTAA

t4-15.nt

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 ATTGAAAACAAGAGCCTGAAAAACAGAAACAAAATGCGAGAAAATAATCCCTACGGTATCAATTCAACGGTAG
 AAATAAGGAATCAAATCAAATCCAAAAGCATTGAGAAGTACTACAAGCAAGCTTATCCGATTCAACATTCA
 TCTTGATTTAGCATCACAAGAGAAAAGGAATTCTAAAACCAGAAGATAAAATCTTGCCCACACAGGGAAAGTG

TABLE 1. Nucleotide and Amino Acid Sequences

GAGTCTTGAGCATCTAATAAATAAAAATTGTTAGACTTTAAAGCCCCAGAAAATCCAAAAGCTCAACTTAA
 AAAATTCAAAGAAATTAAAATATTGAGAATTCTCAAAATCAAGACTTATTATTGTCTAACCTTAAAGA
 TAAAATAACAACAACTATTAAACATCATGCTCAATCCCCAAAGCACATCCAAAACCCAAAGATTATTTA
 AAAGACCTTAAAGACACAATTAAAAGGGTACTGGTGAGAAATACTTAAATCCTATCTATAGATTCAAATAAAAA
 ACAAAAAAGATTATCATTCAATAGATTACAACAAAGTACTATTAGCGAAAAAACAAATAGAATTGGACCTACTGCC
 TCACGAACAAGTCTTCAAATGAATAAAATTCACTAAA

f4-15.aa

MSKKVILILL EILILSCDLS INKEQKTKEK TSEKQESEKQ NIEKQEPEKQ KQNAAKI IPT
 VSIQTVEIRE SNQIPKSIEK YYKQAYPIQT FTLDTSITRE KEFLKPEDKI LPTQGKVESL
 SILINKKLLD FKAPENPKSS TLKNFKEIKN IENFFQNQDL LFVLTLDKN NNNTINIMLN
 PPNDIQKPKD YILKDLKDTI KKGTGEKYLN PIYRFQIKNK KDYHSIDYNTK VTISEKTIEL
 DLLPHEQVFQ MNKNFTKILD TITDLNNLKL VIQKELV

t4-15.aa

CDLSINKEQKTKEKTSEKQESEKQNIEKQEPEKQKQNAAKI IPTVSIQTVEIRESNQIPKSIEKYYKQAYPIQTFT
 LDTSITREKEFLKPEDKILPTQGKVESLSILINKKLLDFKAPENPKSSTLKNFKEIKNENFFQNQDLLFVLTLDK
 KNNNNNTINIMLNPPNDIQKPKDYLKDLKDTIKKGTGEKYLNPIYRFQIKNKKDYHSIDYNTKVTISEKTIELDLLP
 HEQVFQMNKNFTK

f4-50.nt

TAGAAGGAGG AAAAATGAA AATTGGAAAG CTTAAATTCAA TAGTTATAGC CTTGTTTTT
 AACTATTGG TCGCATGTAG TATTGGATTA GTAGAAAGAA CAAATGCAGC TCTTGAATCG
 TCCTCTAAGG ATTTAAAAAA CAAAATTAA AAAATAAAA AAGAAGCCAC GGGAAAAGGT
 GTACTTTTG AAGCTTTAC AGGTCTTAAA ACCGGTTCCA AGGTAACAAG TGGTGGACTA
 GCCTTAAGAG AAGCAAAAGT ACAAGCCATT GTTGAACACAG GAAAGTTCT TAAGATAATA
 GAAGAAGAAG CTTAAAGCT TAAAGAAACT GGAAACAGTG GTCAATTCTT GGCTATGTTT
 GACTTAATGC TTGAGGTTGT AGAATCGCTA GAAGACGTTG GAATAATAGG CTTAAAAGCC
 CGTGTAGGAGGAAATCTAA AAATAATCCT ATAAACACAG CTGAAAGATT GCTTGCAGCT
 AAAGCTCAAATAGAAAATCA ACTTAAAGTG GTTAAGGAAA AACAAAATAT TGAAAATGGT
 GGAGAGAAAA AAAATAATAA AAGCAAAAAA AAGAAATAA

t4-50.nt

ATGTAGTATTGGATTAGTAGAAAGAACAAATGCAGCTCTGAATGTCCTCTAAGGATTAAAAACAAAATTAA
 AAAATAAAAAAGAACGCCACGGAAAAGGTGTACTTTTGAAAGCTTTACAGGTCTAAAACCGGTTCCAAGGTAA
 CAAGTGGTGGACTAGCCTTAAGAGAAGCAAAAGTACAAGCCATTGTTGAAACAGGAAAGTCTTAAGATAATAGA
 AGAAGAACCTTAAAGCTTAAAGAAACTGGAAACAGTGGTCAATTCTGCTATGTTGACTTAATGCTTGAGGTT
 GTAGAATCGCTAGAAGACGTTGGAATAATAGGCTTAAAGCCCGTGTAGAGGAATCTAAAATAATCCTATAA
 ACACAGCTGAAAGATTGCTTGCGGCTAAAGCTCAAATAGAAAATCAACTTAAAGTGGTTAAGGAAAACAAAATAT
 TGAAAATGGTGGAGAGAAAAAATAATAAAGCAAAAAAGAAA

f4-50.aa

KEEKMKIGKL NSIVIALFFK LLVACSIGLV ERTNAALESS SKDLKNKILK IKKEATGKGV
 LFEAFTGLKT GSKVTSGGLA LREAKVQAIV ETGKFLKIIIE EEALKLKETG NSQFLAMFD
 LMLEVVESLE DVGIIGLKAR VLEESKNNPI NTAERLLAAK AQIENQLKVV KEKQNIENGG
 EKKNNKSKKK K

t4-50.aa

TABLE 1. Nucleotide and Amino Acid Sequences

CSIGLVERTNAALESSSKDLKNKILKIKKEATGKGVLFEAFTGLKTGSKVTSGGLALREAKVQAIETGKFLKII
EEALKLKETGNSGQFLAMFDLMLLEVVESLEDVGIIGLKARVLEESKNNPINTAERLLAAKAQIENQLKVVKEKQNI
ENGGEKIQDNKSKKKK

f4-66.nt

TAATTTTAA AATTTAAATA TTTACATAAT AGTAATGTGT GTGGGAGACG TATGAAAAAT
ATTTTATTAT TTGTTATTTT ATTATTCTTT TCTTGTAAAG AATTTAATTAA TTCTGATCTT
AGGAGAAGGC CTTCAAAGGT TTTAAATGCT TCTAATGGTG CATCAAATAA AGAACTTAA
ATTTCTTTG TAGATTCTTT AAATGATGAT CAAAAAGAAG CTTTGTTTT TCTTGAACAG
CTAGTTCTTG ATAGCAATCC CGACAAGTTT AATCAAATT TTAATTAAA TGAAGAGAAG
GTAAGAAA TCCTTGTAC TGTTGTTAAG TGTTAAAGG CCAAAAGAAA GGCTAAAATG
GCTCTTGAGA GCTCAAATGT TGCAAATGTT GCCAATGCTA AACAGCAATT GCTACAGGTT
GAAATTAACCTT ACATAGATAA TTTGCGACAA TCTTTATGA CTACTAAAAA CATTGAAGAG
GCTTGTAAATC TTGTAAAAAA TTATGATGCA TCTGCTTCGT TTTAA

t4-66.nt

TTGTAAGAATTAAATTATTCTGATCTTAGGAGAAGGCCTCAAGGTTTAAATGCTTCTAATGGTCATCAAAT
AAAGAACTTAAATTCTTTGTAGATTCTTAAATGATGATCAAAGAAGCTTGTGTTTCTTGAACAGGTAG
TTCTGATAGCAATCCGACAAGTTAATCAAATTAAATTAAATGAAGAGAAGGTTAAAGAAATGCTTGTAC
TGTTGTTAAGTGTAAAGGCCAAAGAAAGGCTAAATGGCTTGTGAGAGCTCAAATGTTGCAAATGTTGCCAAT
GCTAAACAGCAATTGCTACAGGTTGAAAAACTTACATAGATAATTGCGACAATCTTTATGACTACTAAAACA
TTGAAGAGGCTTGTAACTTGTAAAAATTATGATGCATCTGCTTCGTTT

f4-66.aa

FLKFKYLHNS NVCGRRMKNI LLFVILLFFS CKEFNYSDLR RRPSKVLNAS NGASNKELKI
SFVDSLNDQ KEALFFLEQV VLDSNPDKFN QIFNLNEEKV KEMLVTUVKC LKAKRKAKMA
LESSNVANVA NAKQQLLQVE KTYIDNLRQS FMTTKNIEEA CNLVKNYDAS ASF

t4-66.aa

CKEFNYSDLRRRPSKVLNASNGASNKELKISFVDSLNDQKEALFFLEQVVLDSNPDKFNQIFNLNEEKVKEMLVT
VVKCLKAKRKAQMALESSNVANVANAKQQLLQVEKTYIDNLRQSFMTTKNIEEACNLVKNYDASASF

f42-1.nt

TAATTTTAA AATCTAAGGA GAAGAGATT ATGAACAAAA AATTTCTAT TTCATTATTA
TCTACAATAT TAGCCTTCTT GTTAGTATTA GGTGTTGATT TGTCAAGCAA TAATGCTGAA
AACAAAATGG ATGATATTTT TAATTTAGAA AAGAAATACA TGGATAATTCA AAATTATAAA
TGTTAAGTA AAAATGAGGC TATAGTTAA AATTCTAAAA TTAAATTAGG TGTAAATAAT
ACTAGAAGTC GTTCTTATTTC TTCTAGAGAG ACTAATGTTT CGGATTCCCTA TAATAAAACC
TATTCATATT GCAAAGCAA CTGA

t42-1.nt

TTGTGATTTGTCAAGCAATAATGCTGAAAACAAAATGGATGATATTTTAATTAGAAAAGAAATACATGGATAAT
TCAATTATAATGTTAAGTAAAATGAGGCTATAGTTAAAATCTAAATTAAATTAGGTGTAATAACTA
GAAGTCGTTCTTATTCTTAGAGAGACTAATGTTCGGATTCCCTATAATAAAACCTATTCAATTGCAAAAGCAA
C

f42-1.aa

LLKSKEKRFM NKKFSISLLS TILAFLLVLG CDLSSNNAEN KMDDIFNLEK KYMDNSNYKC
LSKNEAIVKN SKIKLGVNNT RSRSYSSRET NVSDSYNKT SYCKSN

TABLE 1. Nucleotide and Amino Acid Sequences

t42-1.aa

CDLSSNNAENKMDIFNLEKKYMDNSNYKLSKNEAIVKNSKIKLGVNNTRSRSYSSRETNVSDSYNKTYSYCKSN

f43-3.nt

TGAATATTA TAATAAAAAA AGGAATAANA ATGAAAATTA TCAACATATT ATTTTGTAA
 TTTTTACTAA TGCTAACAG CTGTAATTCT AATGATACTA ATACTAGCCA AACAAAAAGT
 AGACAAAAAC GTGATTTAAC CCAAAAAGAA GCAACACAAG AAAAACCAAA ATCTAAAGAA
 GACCTGCTA GAGAAAAGCT ATCTGAAGAC CAAAAAACAC ATCTTGACTG GTTAAAAACC
 GCTTAACTG GTGCTGGAGA ATTTGATAAA TTTTAGGAT ATGACGAAGA CAAAATAAAA
 GGTGCACTTA ATCATATAAA GAGTGAACCT GATAAGTGT A CTGGGATAA TTCTGAACAA
 CAAAAAGCA CCTTCAAAGA GGTGGTTAAG GGGGCTTGT GTGGCGGTAT AGATAGTTT
 GCAACTAGTG CAAGTAGTAC CTGCCAACGCT CAGCAATAA

t43-3.nt

CTGTAATTCTAATGATACTAATAGCCAAACAAAAAGTAGACACAAAACGTGATTTAACCCAAAAGAACAGCAACA
 CAAGAAAAACCAAAATCTAAAGAAGACCTGCTTAGAGAAAAGCTATCTGAAGACCAAAACACATCTGACTGGT
 TAAAAACCGCTTAACTGGTGTGGAGAATTGATAAAATTAGGATATGACGAAGACAAAATAAAAGGTGCACT
 TAATCATATAAAAGAGTGAACCTGATAAGTGTACTGGGATAATTCTGAACACAAAAAGCACCTTCAAAGAGGTG
 GTTAAGGGGGCTTGGTGGCGGTATAGATAGTTTGCAACTAGTGCAAGTAGTACCTGCCAACGCTCAGCAA

f43-3.aa

ILIIKKGIXM KIINILFCLF LLMLNSCNSN DTNTSQTKSR QRDLTQKEA TQEKPKSSED
 LLREKLSEDQ KTHLDWLKTA LTGAGEFDKF LGYDEDKIKG ALNIKSELD KCTGDNSEQQ
 KSTFKEVVKG ALGGGIDSFA TSASSTCQAQ Q

t43-3.aa

CNSNDTNTSQTKSRQKRDLTQKEATQEKPKSSEDLLREKLSEDQKTHLDWLKTALTGAGEFDKFLGYDEDKIKGAL
 NHIKSELDKCTGDNSEQQKSTFKEVVKGALGGGIDSFATSASSTCQAQ

f45-2.nt

TAGGAGAGAA TAATTATGAA TAAAAAAACA TTGATTATTT GTGCTGTTT TCCGCTGATA
 ATTTCTTGCA AGAATTTGCA AACTGGTAA GATATAAAAC AAAATTCAGA AGGGAAAATT
 AAAGGATTG TAAATAAGAT TTTAGATCCA GTAAAGGATA AAATTGCTTC AAGTGGTACA
 AAAGTAGATG AAGTAGCAA AAAATTACAA GAAGAAGAAA AAGAAGAATT AATGCAGGGC
 GATGATCCTA ATGGCAGTGG AATAAATCCG CCACCACTAT TGCCGAAAA TATTCAACAT
 AATGCATTAG TATTAAAGC AATAGAACAA AGTGTGGTC AACAAAGAAAA AAAAGTAGAA
 GAAGCTGAAG CTAAGTTGA AGAAAATAAA GAAAAACAAG AGAATACAGA AGAAAACATT
 AAAGAAAAAG AAATAATAGA CGAACAAAC AAACAAGAAT TAGCTAAAGC TAAAGAAGAA
 GAACAACAAA AAGAACAAAA AAGACATCAA GAAGAGCAAC AAAGAAAAGC TAAAGCAGAA
 AAAGAAAAAA GAGAAAGAGA AGAGGCAGAA CAACAAAAAC GACAACAAGA AGAGGAAGAA
 AAAAGGCAAG TTGATAACCA AATTAAAACA CTTATAGCTA AAATAGATGA GATCAATGAA
 AATATTGATG TTATAAAATG GCAAACGACT GTAGGCCAC AAGGCGTTAT AGATAGAATT
 ACTGGGCTG TGTATGATGA TTTTACCAAT GGCAATAATT CTATACCGCA AACTTGGGAG
 GGGTTAGAAG AGGAATCAGA AGACGAAGGA TTAGGAAAAT TATTGAAAGA ATTGACTGAT
 GCTAGGGACG CGCTAAGAAC TAAATTAAAT GAAGGCAATA AACCATATAC TGGTTACGAA
 GAGCCTAAGT TAAAAGAAAG TGTAATGTT AGCGAAATTA AAGAAGATTT AGAAAAAATT
 AAATCAAAAT TAGAAGAAGT TAAAAAATAT CTTAAAGATA GTTCTAAATT TGAAGAATT
 AAAGGATACA TCAGTGACAG TCAGTAA

t45-2.nt

TABLE 1. Nucleotide and Amino Acid Sequences

TTGCAAGAATTTGCAACTGGTAAAGATATAAACAAATTCAAGAAGGGAAAATTAAAGGATTTGTAATAAGATT
 TTAGATCCAGTAAAGGATAAAATTGCTTCAGTGGTACAAAAGTAGATGAAGTAGCAAAAAATTACAAGAAGAAG
 AAAAGAAGAATTAATGCAGGGCGATGATCCTAATGGCAGTGGATAAAATCCGCCACCAAGTATTGCCGAAAATAT
 TCACAATAATGCATTAGTATTAAAGCAATAGAACAAAGTGTAGGGTCAACAAGAAAAAAAGTAGAAGAAGCTGAA
 GCTAAAGTTGAAGAAAATAAAGAAAAACAGAGAACATACAGAACAGAAACATAAAGAAAAGAAATAATAGACGAAC
 AAAACAAACAGAATTAGCTAAAGAACAGAACAAAAAGAACACAAAAAGACATCAAGAACAGAGCAACA
 AAGAAAAGCTAAAGCAGAAAAAGAAAAAGAGAACAGAGAACAGAGGCAAGAACACAAAAACGACAACAAGAAGAGGAA
 GAAAAAAGGCAAGTGTAGAACCAATTAAACACTTATAGCTAAATAGATGAGATCAATGAAAATATTGATGTTA
 TAAAATGGCAAACGACTGTAGGCCACAAGGCGTTAGATAGAACATTACTGGCCTGTGTATGATGATTTACCAA
 TGGCAATAATTCTATACCGCAGACTTGGGAGGGTTAGAAGAGAACATCAGAACAGAACAGGATTAGGAAAATTATTG
 AAAGAATTGAGTGTAGCTAGGGACGCGCTAGAACATTAAATGAAGGCAATAACCATATACTGGTTACGAAG
 AGCCTAAGTAAAAGAAAAGTGTAAATGTTAGCGAAATTAAAGAACAGATTAGAAAATTTAAACAAATTAGAAGA
 AGTAAAAAATATCTTAAAGATAGTTCTAAATTGAGAACATTAAAGGATACATCAGTACAGTCAG

f45-2.aa

ERIIMNKCTL IICAVFALII SCKNFATGKD IKQNSEGKIK GFVNKILDGV KDKIASSGK
 VDEVAKKLQE EKEELMQGD DPNGSGINPP PVL PENIHNN ALVLKAIIEQS DGQQEKKV
 AEAKVEENKE KQENTEENIK EKEIIDEQNQ QELAKAKEEE QQKEQKRHQE EQQRKAKAEK
 EKREREEAQ QKRQEEEEK RQVDNQIKTL IAKIDEINEN IDVIKWQTTV GPQGVIDRIT
 GPVYDDFTNG NNSIRETWEGL EEESEDEGL GKLLKELSDA RDALRTKLNE GNKPYTGYEE
 PKLKESYNVS EIKEDLEKLK SKLEEVKKYL KDSSKFEIYK GYISDSQ

t45-2.aa

CKNFATGKDIKQNSEGKIKGFVNKILDGVKDKIASSGKVDEVAKKLQEEEKEELMQGDDPNGSGINPPPVL PENI
 HNNALVLKAIIEQSDGQKEKKVVEEAKEVENKEKQENTEENIKEKEIIDEQNQKQELAKAKEEEQQKEQKRHQEEQQ
 RAKAKAEKEKREREEAEQQKRQEEEEKRQVDNQIKTLIAKIDEINENIDVIKWQTTVGPQGVIDRITGPVYDDFTN
 GNNSIRETWEGL EEESEDEGLKLLKELSDARDALRTKLNEGNKPYTGYEEPKLKESVNVEIKA
 VKKYLKDSSKFEIKGYISDSQ

f47-2.nt

TGAATATTAA TAATAAAAAA AGGAGTAACA ATGAAAATCA TCAACATATT ATTTTGTATA
 TCTTTGCTAC TACTAAATAG CTGTAATTCC AATGATAATG ACACCTTAAA AAACAATGCC
 CAACAAACAA AAAGCAGGAA AAAACGTGAT TTAAGCCAAG AAGAACTGCC ACAACAAGAA
 AAAATCACTT TAACATCCGA CGAAGAAAA ATGTTTACTT CATTAAATCAA TGTGTTAAA
 TACACAAATTG AAAAATTAAA CAATGAAATA CAAGGGTGCA TGAATGGAAA CAAAGTAAA
 TGTAAATGACT TCTTGATG GCTTTCTGAA GATAATTCAA AACAAAAGA ATTAGCTGGT
 GCTTTTACCA AGGTTTACAA CTTCTTAAA TCAAAAGCAC AAAATGAAAC TTTTGATACT
 TATATTAAAG GAGCTATTGA TTGAAAAAA AACACTCCAC AAGATTGTAA TAAAATAAT
 GAAATATGGG GAGGTGGACA ACTTANTAGN GCAATATTAG AG

t47-2.nt

CTGTAATTCCAATGATAATGACACTTTAAAAACAAATGCCAACAAACAAAAGCAGGAAAAACGTGATTTAAC
 CAAGAAGAACTGCCAACAAAGAAAAATCACTTAAACATCCGACGAAGAAAAATGTTACTCATTAAATCAATG
 TGTTAAATACACAATTGAAAAATTAAACATGAAATACAAGGGTGCATGAATGAAACAAAGTAAATGAA
 CTTCTTGATTGGCTTCTGAAAGATATT
 CAAAACAAAAGAATTAGCTGGTCTTTACCAAGGTTACAACCTTAAAATCAAAAGCACAAAATGAAAC
 TTGATACTTATATTAAAGGAGCTATTGATGTTAAAAACACTCCACAAAGATTGTAAATAAAATAATGAA

f47-2.aa

ILIIKKGVMT KIINILFCIS LLLLNSCNSN DNDTLKNNAQ QTCSRKKRDL SQEELPQQEK

TABLE 1. Nucleotide and Amino Acid Sequences

ITLTSDEEKM FTSLINVFKY TIEKLNNEIQ GCMNGNKSCK NDFFDWLS ED IQKQKELAGA
FTKVYNFLKS KAQN EFTDY IKGAIDCKKN TPQDCNKNNE IWGGQLXXA IF

t47-2.aa

CNSNDNDTLKNNAQQTCSRKKRDLSQEELPQQEKITLTSDEEKMFTSLINVFKYTIEKLNNEIQGCMNGNKSCKND
FFDWLSEDIQKQKELAGAFTKVYNFLKSQAQN EFTDYIKGAIDCKNTPQDCNKNNE

f49-2.nt

TAAATGTTCA AAACAATCAT TAAACAAAAA AATATGAAAA AAATTCAG TGCAATTAA
TTAACAACTT TCTTGT TTTT TATTAATTGT AAAAGCCAAG TTGCTGATAA GGCAGTG
ACGGGGATTG CTAAGGGAAT AAAGGAGATT GTTGAAGCTG CTGGGGGAG TGAAAAGCTG
AAAGTTGCTG CTGCTGAAGG GGAGAATAAT GAAAAGGAG GGAAGTTGTT TGGGAAGGCT
GGTGCTGGTA ATGCTGGGA CAGTGAGGCT GCTAGCAAGG CGGCTGGTGC TGTTAGTGT
GTTAGTGGGG AGCAGATATT AAGTGCAGATT GTTAAGGCTG CTGGTCAAGG TGCGCAGGAT
GGAGAGAAGC CTGGGGAGGC TAAAATCCG ATTGCTGCTG CTATTGGAA CGGTAAATGAG
GATGGTCCGG AGTTAAGGA TGAGATGAAG AAGGATGATC AGATTGCTGC TGCTATTGCT
TTGAGGGGGG TGGCTAAGGA TGGAAAGTTT GCTGTGAAGA ATGATGAGAA AGGGAAAGGCT
GAGGGGGCTA TTAAGGGAGC TGGCGAGTTG TTGGATAAGC TGGTAAAAGC TGTAAAGACA
GCTGAGGGGG CTTCAAGTGG TACTGCTGCA ATTGGAGAAG TTGTGGCTGA TGATAATGCT
GCGAAGGTTG CTGATAAGGC GAGTGTGAAG GGGATTGCTA AGGGATAAA GGAGATTGTT
GAAGCTGCTG GGGGGAGTAA AAAGCTGAA GTTGCCTGCTG CTAAAGAGGG CAATGAAAAG
GCAGGGAAAGT TGTTGGAA AGTTGATGCT GCTCATGCTG GGGACAGTGA GGCTGCTAGC
AAGGCGGCTG GTGCTGTTAG TGCTGTTAGT GGGGAGCAGA TATTAAGTGC GATTGTTAAG
GCTGCTGGTG CGGCTGCTGG TGATCAGGAG GGAAAGAACG CTGGGATGCA TAAAATCCG
ATTGCTGCTG CTATTGGAA GGGTGTGCG GAGAATGGTG CGGAGTTAA TCATGATGGG
ATGAAGAAGG ATGATCAGAT TGCTGCTGCT ATTGCTTGA GGGGGATGGC TAAGGATGGA
AAGTTGCTG TGAAGAGTGG TGGTGGTGAG AAAGGGAGG CTGAGGGGGC TATTAAGGGA
GCTGCTGAGT TGTTGGATAA GCTGGTAAA GCTGTAAAGA CAGCTGAGGG GGCTTCAAGT
GGTACTGATG CAATTGGAGA AGTTGTTGCT AATGCTGCTG CTGCAAAGGT TGCTGATAAG
GCGAGTGTGA CGGGGATTGC TAAGGGATA AAGGAGATTG TTGAAGCTGC TGGGGGGAGT
GAAAAGCTGA AAGTTGCTGC TGCTACAGGG GAGAGTATA AAGGGGCAGG GAAGTTGTT
GGGAAGGCTG GTGCTGGTG TAATGCTGGG GACAGTGGAGG CTGCTAGCAA GGCGGCTGGT
GCTGTTAGTG CTGTTAGTGG GGAGCAGATA TTAAGTGCAG TTGTTAAGGC TGCTGATGCG
GCTGATCAGG AGGGAAAGAA GCCTGGGAT GCTANAAATC CGATTGCTGC TGCTATTGGG
AAGGGTNATG NGGAGAATGG TGCGGAGTTT AANNATGANG GATGA

t49-2.nt

TTGTAAGGCAAGTTGCTGATAAGGCGAGTGTGACGGGATTGCTAAGGAATAAGGAGATGTTGAAGCTGCT
GGGGGGAGTGAAGCTGAAAGTTGCTGCTGCTGAAGGGAGATAATGAAAAGGCAAGGAAGTTGTTGGGAAGG
CTGGTCTGGTAATGCTGGGACAGTGAGGCTGCTAGCAAGGCGCTGGTGTAGTGTGCTGTTAGTGGGAGCA
GATATTAAAGTGCATTGTTAAGGCTGCTGGTGAGGCTGCGCAGGATGGAGAGAAGCCTGGGGAGGCTAAAATCCG
ATTGCTGCTGCTATTGGAAACGGTAATGAGGATGGTGCAGGTTAAGGATGAGATGAAGAAGGATGATCAGATTG
CTGCTGCTATTGCTTTGAGGGGGATGGCTAAGGATGGAAAGTTGCTGTGAAGAATGATGAGAAAGGAAAGGCTGA
GGGGGCTATTAAG

f49-2.aa

MFKTIIKQKN MKKISSAILL TTFVFINCK SQVADKASVT GIAKGKEIV EAAGGSEKLK
VAAAEGENNE KAGKLFKGAG AGNAGDSEAA SKAAGAVSAV SGEQILSAIV KAAGEAAQDG
EKPGEAKNPI AAAIGKGNED GAEFKDEMKK DDQIAAAIAL RGMAKDGKFA VKNDEKGKAE
GAIKGAGELL DKLVKAVKTA EGASSGTAAI GEVVAADDNAA KVADKASVKG IAKGKEIVE
AAGGSKKLV AAAKEGNEKA GKLFGKVDAA HAGDSEASK AAGAVSAVSG EQILSAIVKA
AGAAAGDQEG KKPQDAKNPI AAAIGKGDAE NGAEFNHDGM KKDDQIAAAI ALRGMAKDGGK

TABLE 1. Nucleotide and Amino Acid Sequences

FAVKSGGGEK GKAEGAIKGA AELLDKLVKA VKTAEGASSG TDAIGEVVAN AGAAKVADKA
 SVTGIAKGIK EIVEAAGGSE KLKVAAATGE SNKGAGKLFG KAGAGANAGD SEAASKAAGA
 VSAVSQEQL SAIVKAADAA DQEKKPGDA XNPIAAAGK GXXENGAEX XXG

t49-2.aa

CKSQVADKASVTGIAKGIKEIVEAAGGSEKLVAAAEGENNEKAGKLFGKAGAGNAGDSEAASKAAGAVSAVSGEQ
 ILSAIVKAAGEAAQDGEPGEAKNPIAAAGKGNEDGAEFKDEMKKDDQIAAAIALRGMAKDGFVKNDEKGKAE
 GAIK

f5-14.nt

TAGAAATTCA AAACAAAGGA GAAAACAAAA AGTATGAATA AAAAAATATT GATTATTTT
 GCTGTTTTG CACTTATAAT TTCTTGAAA AATTATGCAA CTGGTAAAGA TATAAAACAA
 AATGCAAAAG GGAAAATTAA AGGATTTTA GATAAGGTT TAGATCCAGC AAAAGATAAA
 ATTACTTCAA GTACTTCAA AGTAGATGAA TTGCAAAAA AATTACAAGA AGAAGATGAA
 GATAATGAAT TAATGCAGGG CGATGATCCT AATAACAGAG CAATAGCACT GTTACCAAGTA
 TTGCCGAAA ATAGTCATGA CAATCCACCA GTACCAAAAG TAAAAGCAGC AGCACAAAGT
 GGTGGTCAAC AAGAAGACCA AAAAGCAAA GAATCTAAAG ATAAAGTGA CGAAGAAAAA
 GAAGTTGTAG AGGAGAAAAA AGAAGAACAA GATAGTAAA AAGAAAAAGT GGAGAAGCAA
 AGTCAAAAGC AAAAGAAGA AGAGAGAAC TCTAAAGAAG AACACAAAA ACAAGAAGAA
 GCAAAAGCTA GAGCAGATAG AGAAAGAGAA GAACGACTAA AACACAAAGA ACAAAAAAGA
 CAACAGGAAG AAGCTAGGGT TAAAGCAGAA AAAGAAAAAC AAGAAAGAGA GGAACAACAA
 AAACAAGAAG AAGAAAAGAA AGTTAAATAT AAAATTAAA CACTTACAGA CAAAATAGAT
 GAAATAAATA AGGATATTGA TGGTATAAAT GGTAACACAA TTGTAGGAGC AGAAGAAGTT
 ATAGATAAAA TTACGGGCC TGTATATGAT GATTTACTG ATGGGAATAA AGCTATATAC
 AAAACTTGGG GAGATTAGA GGATGAAGAA GGCGAAGAAT TAGGAAAATT ATTGAAAGAA
 TTGAGTGATA CTAGACATAA TTTAAGAACC AAATTAAATG AGGGTAATAA AGCATATATT
 GTTCTAGAAA AGGAGCCTAA TTTAAAGAA AATGTAATG TTAGTGATAT TCAATCAGAT
 TTAGAAAAAT TAAAATCAGG ATTAGAAGAA GTTAAAAAAT ATTTGAAAA TGAAGATAAT
 TTTGAAGAAA TTAAAGGATA CATTGAGGAT AGTAATTCAAT ATTGA

t5-14.nt

TTGTAAAAATTATGCAACTGGTAAAGATATAAAACAAAATGCAAAAGGGAAAATTAAAGGATTTAGATAAGGTT
 TTAGATCCAGCAAAAGATAAAATTACTTCAGTGTCAAAAGTAGATGAATTAGCAAAAAATTACAAGAAGAAG
 ATGAAGATAATGAATAATGCAGGGCGATGATCCTATAACAGAGCAATAGCACTGTTACCAAGTATTGCCGAAA
 TAGTCATGACAATCCACCACTACCAAAAGTAAAGCAGCAGCACAAAGTGGTGGTCAACAAGAAGACCAAAAGCA
 AAAGAATCTAAAGATAAAAGTTGAGGAAGAAAAGAAGTTGTAGAGGAGAAAAAGAAGAACAAAGATAGTAAAAAG
 AAAAGTGGAGAACAGTCAAAAGCAAAAGAACAGAGAGAGAACACTCTAAAGAACAAACAAAACAAGAAGA
 AGCAAAAGCTAGAGCAGATAGAGAACAGACTAAAACAACAAGAACAAAAAGAACACAGGAAGAACAGCT
 AGGGTTAACAGAAAAAGAAAACAAGAACAGAGAGAACACAAGAACAGAACAGAACAGAACAGCTAAATATA
 AAATTAAAACACTTACAGACAAATAGATGAAATAATAAGGATATTGATGGTATAATGGTAAAACAATTGTAGG
 AGCAGAACAGTTATAGATAAAATTACGGGCCTGTATATGATGATTTACTGATGGAATAAAGCTATATACAAA
 ACTTGGGAGATTTAGAGGATGAAGAACGGCAAGAATTAGGAAAATTATTGAAAGAATTGAGTGATACTAGACATA
 ATTTAAGAACCAATTAAATGAGGGTAATAAAGCATATATTGTTCTAGAAAAGGAACCTAATTAAAAGAAAATGT
 AAATGTTAGTGATATTCAATCAGATTAGAAAAATTAAATCAGGATTAGAACAGTTAAAATATTGAAAAT
 GAAGATAATTGAAAGAAATTAAAGGATACATTGAGGATAGTAATTCAAT

f5-14.aa

KFKTKEKTKS MNKKILIIIFA VFALIISCKN YATGKDIKQN AKGKIKGFLD KVLDPAKDKI
 TSSSSKVDEL AKKLQEEDED NELMQGDDPN NRAIALPVL PENSHDNPPV PKVKAQQSG
 GQQEDQKAKE SKDKVEEKE VVEEKKEEQD SKKEKVEKQS QKQKEEERNS KEEQQKQEEA
 KARADRREEE RLKQQEOKRQ QEEARVKAEK EKQEREQQK QEEEKVKYK IKTLDKIDE
 INKDIDGING KTIVGAEVVI DKITGPVYDD FTDGNKAIYK TWGDLDEEG EELGKLLKEL

TABLE 1. Nucleotide and Amino Acid Sequences

SDTRHNLRTK LNEGNKAYIV LEKEPNLKEN VNVSIDIQSDL EKLKSGLEEV KKYFENEDNF
EEIKGYIEDS NSY

t5-14.aa

CKNYATGKDIKQNAKGKIKGFLDKVLDPAKDKITSSSSKVDELAKKLQEEDEDNELMQGDDPNNRAIALLPVL PEN
SHDNPPVKVAAAQSQQQEDQKAKESDKVEEEKEVVEEKKEEQDSKKEKVEKQSQKQKEEERNSEEQQKQEE
AKARADREREERLKQQEQKRQQEEARVKAKEKEQEREEQQKQEEKKVKYKIKTLTDKIDEINKDIDGINGKTI VG
AEEVIDKITGPVYDDFTDGNKAIYKTWGDLEDEEGLKLLKELSDTRHNLRTKLNEGNKAYIVLEKEPNLKEN V
NVSDIQSLEKLKSGLEEVKKYFENEDNFEEIKGYIEDNSY

f5-15.nt

TAACCTTATGA ATAAGAAAAT GAAAATGTTT ATTATTTGTG CTGTTTTGC ATTGATGATT
TCTTGCAAGA ATTATGCAAG TGGTGAAAAT CTAAAAAATT CAGAACAAAAA TCTAGAAAGT
TCAGAACAAA ATGTAaaaaAA AACAGAACAA GAGATAaaaaAA ACAAGTTGA AGGATTTTA
GAAATTCTAG AGACAAAAGA TTTATCTAAA TTAGATGAAA AAGATACAAA AGAAATTGAA
AAACAAATTG AAGAATTAAA GAATAAAAATA GAAAATTAG ATTCTAAAAA AACTTCTATT
GAAACATATT CTGAGTATGA AGAAAAAATA AACAAAATAA AAGAAAATT GAAAGGAAAAA
GGACTTGAAG ATAAATTAA GGAGCTTGAAG GAGAGTTAG CAAAGAAAAA GGGGGAGAGA
AAAAAAGCTT TACAAGAGGC CAAACAGAAA TTTGAAGAAT ATAAAAAACA AGTAGATACT
TCAACTGGGA AAACTCAAGG CGACAGGTCT AAAAACCGAG GTGGTGTGAGTCAAGGACTT
TGGCAGTGTG CCAATGAATT AGGTTGGGT GTAAGTTATT CTAATGGCGG CAGTGACAAC
AGCAAACTTG ATGAATTAGC AAACAAAGTT ATAGATGATT CTCTTAAAAA GATTGAAGAA
GAACCTTAAGG GAATAGAAGA AGATAaaaaAA GAATAA

t5-15.nt

TTGCAAGAATTATGCAAGTGGTAAAATCTAAAAAATTCTAGAACAAAATCTAGAAAGTTCTAGAACAAAATGTA AAAA
AAAACAGAACAAAGAGATAAAAAAAACAAGTTGAAGGATTTAGAAATTCTAGAGACAAAAGATTCTAAATTAG
ATGAAAAGATAACAAAAGAAATTGAAAACAAATTCTAGAACAAATTAAAGAATAAAATAGAAAATTAGATTCTAAAAA
AACTTCTATTGAAACATATTCTGAGTATGAAGAAAAATAAACAAAATAAAGAAAATTGAAAGGAAAAGGACTT
GAAGATAAATTAAAGAGCTTGAAGAGAGTTAGCTAACAGAAAAGGGGGAGAGAAAAAAAGCTTACAAGAGGCCA
AACAGAAATTGAAAGAATATAAAAACAAGTAGATACTTCAACTGGAAAACCTCAAGGCACAGGTCTAAAACCG
AGGTGGTGTGGAGTGCAAGCTGGCAGTGTGCCATTGAATTAGTTGGGTGTAAGTTATTCTAATGGCGGAGT
GACAACAGCAAACTGATGAATTAGCTAACAAAGTTATAGATGATTCTTAAAAGATTGAAGAAACTTAAGG
GAATAGAAGAAGATAAAAAGAA

f5-15.aa

LMNKKMFMFI ICAVFALMIS CKNYASGENL KNSEQNLESS EQNVKKTEQE IKKQVEGFL E
ILETKDLSKL DEKDTKEIEK QIQLKNKIE KLDKKSIE TYSEYEKIN KIKEKLKGKG
LEDKFKELEE SLAKKKGERK KALQEAKQKF EYKQVDT TGKTQGDRSK NRGGVGVQAW
QCANELGLGV SYSNGGSDNS NTDELANKVI DDSLKKIEEE LKGIEEDKKE

t5-15.aa

CKNYASGENLKNSEQNLESSEQNVKKTEQEIKKQVEGFL EILETKDLSKLDEKDTKEIEKQIQLKNKIEKLDSSKK
TSIETYSEYEKINKIKEKLKGKGLEDKFKELEESLAKKKGERKKALQEAKQFEEYKKQVDTSTGKTQGDRSKNR
GGVGVQAWQCANELGLGVSYSNGGSDNSNTDELANKVIDDSLKKIEEELKGIEEDKKE

f51-2.nt

TAATTGTTTG GGGTTGTGGT AAACTTAAGG CTTATGGAGT GGATTATGAA TAAAAAAATG
AAAATATTAA TTATTTGTGC TGTATTTGTG CTGATAAGTT CTTGCAAGAT TGATGCAACT
GGTAAAGATG CAACTGGTAA AGATGCAACT GGTAAGATG CAACTGGTAA AGATGCAACT
GGTAAAGATG CAGAACAAAA TATAAAACGG AAAGTTCAAG GATTTTGAAG AAAGATTAA

TABLE 1. Nucleotide and Amino Acid Sequences

GATCCAGTAA AGGATAAAAT TGCTTCAAAT GGTCCAATAG CAGATGAATT GGCAAAAAAA
 TTACAAGAAG AAGAAAAGGT AAATAACCGG GAAGAAGAAA ATGATAAAAGC TGTCTTTTA
 GGAGAAGAAT CAAAAGAGGA TGAAGAAGAA AATGAGCAAG CTGTTAATTT AGAAGAAAAA
 AATGCGGAAG AGGATAAGAA AGTTGTTAAT TTAGAAGAGA AAGAATTAGA AGTTAAAAAA
 GAGACTGAAG AAGATGAAGA TAAAGAAGAA ATAGAGAAAC AAAAACAAAGA AGTGGAAAAA
 GCACAAGAAA GAAAACAACG ACAAGAAGAA AAGAAACGAA AAAAACAAAGA ACAGCAAGAA
 GAAAAGAAAAC GAAAACGACA AGAACAAAGA AAAGAAAGGA GAGCTAAAAA CAAAATTAAA
 AAACTTGCAG ATAAAATAGA TGAGATAAGT TGGAATATTG ATGGTATAGA AAGTCAAACA
 AGTGTAAAAC CGAAAGCAGT TATAGATAAA ATTACGGGGC CTGTATATGA TTATTTTACC
 GATGACAACA AAAAAGCTAT ATATAAAACA TGGGGAGATT TAGAAGATGA AGAAGGCAGA
 GGATTGGGAA AATTATTGAA AGAATTGAGT GATACTAGAG ATGAGTTAAG AACCAAATTA
 AATAAAAGATA ATAAAAAATA TTATGCCAT GAAAATGAGC CTCCTCTAAA AGAAAATGTA
 GATGTCAGCG AAATTAAAGA AGATTTAGAA AAAGTAAAAT CAGGATTAGA AAAGGTTAAA
 GAATATCTTA AAGACAATTG TAAATTGAA GAAATTAAAG GATACATCAG TTACAGTCAG
 TAA

t51-2.nt

TTGCAAGATTGATGCAACTGGTAAAGATGCAACTGGTAAAGATGCAACTGGTAAAGATGCAACTGGTAAAGATGCA
 ACTGGTAAAAATGCAGAACAAAATATAAAAGGGAAAGTCAAGGATTTAGAAAAGATTTAGATCCAGTAAAGG
 ATAAAATTGCTTCAAATGGTCAAAGCAGATGAATTGGCAAAAAAATTACAAGAAGAAGAAAAGTAAATAACGG
 GGAAGAAGAAAATGATAAGCTGTCTTTAGGAGAAGAATCAAAGAGGATGAAGAAGAAAATGAGCAAGCTGTT
 AATTAGAAGAAAAAATGCGGAAGAGGATAAGAAGTGTAAATTAGAAGAGAAAATTAGAAGTTAAAAAG
 AGACTGAAGAAGATGAAGATAAGAAGAAATAGAGAAACAAAACAAGAAGTGGAAAAAGCACAAGAAAGAAAACA
 ACGACAAGAAGAAAAGAAACGAAAAAACAAAGAACAGCAAGAAGAAAAGAAACGACAAGAACAAAGAAA
 GAAAGGAGAGCTAAACAAAATTAAAAACTTGCAGTAAATTAGATGAGATAAGTGGAAATTGATGTTAG
 AAAGTCAAACAAGTGTAAAACCGAAAGCAGTTATAGATAAAATTACGGGGCTGTATATGATTATTTACCGATGA
 CAACAAAAAGCTATATATAAAACATGGGGAGATTAGAAGATGAAGAAGCGAAGGATTGGAAAATTATTGAAA
 GAATTGAGTGTAACTAGAGATGAGTTAGAACCAAATTAAAGATAATAAAATATTATGCCATGAAAATG
 AGCCTCCTCTAAAAGAAAATGTAGATGTCAGCGAAATTAAAGAAGATTAGAAAATCAGGATTAGAAAA
 GTTAAAGAATATCTTAAAGACAATTCTAAATTGAAGAAATTAAAGGATACATCAGTTACAGTCAG

f51-2.aa

LFGVVNLRL MEWIMNKKMK IFIIICAVFVL ISSCKIDATG KDATGKDATG KDATGKDATG
 KNAEQNIKGK VQGFLEKILD PVKDKIASNG PIADELAKKL QEEEKVNNGE EENDKAVFLG
 EESKEDEEEN EQAVNLEEKN AEEDKKVVNL EKELEVKEE TEDEDKKEE EKQKQEVEKA
 QERKQRQEEK KRKKQEQQEE KKRKRQEQRK ERRAKNKIKK LADKIDEISW NIDGIESQTS
 VKPKAVIDKI TGPVYDYFTD DNKKAIYKTW GDLEDEEGEG LGKLLKELSD TRDELRTKLN
 KDNKKYYAHE NEPPLKENVD VSEIKEDLEK VKSGLEKVKE YLKDNSKFEE IKGYISYSQ

t51-2.aa

CKIDATGKDATGKDATGKDATGKNAEQNIKGKVQGFLEKILD PVKDKIASNG PIADELAKKL QEEEKVNNNG
 EENDKAVFLG EESKEDEEENEQAVNLEEKNAEEDKKVVNL EKELEVKEE TEDEDKKEE EKQKQEVEKA
 QERKQRKKQEQQEEKKRKRQEQRKERRAKNKIKK LADKIDEISW NIDGIESQTS VPKPAVIDKIT
 NKKAIYKTWDLEDEEGEG LGKLLKELSD TRDELRTKLN KDNKKYYAHENE
 PPLKENVD VSEIKEDLEK VKSGLEK VKEYLKDNSKFEE IKGYISYSQ

f6-21.nt

TAGGCAAAAT TTAAATTAT AAAAACTGT AAGGATGCTT GTATGAAAAT ATTGATAAAA
 AAGTTAAAAG TTGTATTATT TCTCAATTAA ATTAACTTA TTTCTTGTGT TAATGAAAGT
 AATAGAAACA AATTGGTTTT TAAGCTAAAT ATTGGAAGTG AGCCTGCTAC TTTAGATGCT
 CAATTAATAA ACGATACGGT TGGATCAGGG ATTGTAAGCC AAATGTTCT TGGCATTAA
 GATGGAGATC CCAGGACTGG AGGATACAGA CGGGACTTG CTAAAGTTG GGATATTCT

TABLE 1. Nucleotide and Amino Acid Sequences

GATGACGGAG TAGTTTATAC GTTTCATTTA AGAGATAATC TTGTTGGAG TGATGGAGTT
TCCATTACTG CCGAAGAATA A

t6-21.nt

TTGTGTTAATGAAAGTAATAGAACAAATTGGTTTAAGCTAAATATTGGAAGTGAGCCTGCTACTTAGATGCT
CAATTAATAAACGATAACGGTTGGATCAGGGATTGTAAGCCAAATGTTCTGGCATTTAGATGGAGATCCCAGGA
CTGGAGGATACAGACCGGGACTTGCTAAAAGTTGGGATATTCTGATGACGGAGTAGTTATACGTTCATTTAAG
AGATAATCTGTTGGAGTGTGGAGTTCCATTACTGCCGAAGAA

f6-21.aa

AKFKFIKTCK DACMKILIKK LKVVLFLNLI LLISCVNESN RNKLVFKLNI GSEPATLDAQ
LINDTVSGSI VSQMFLGILD GDPRTGGYRP GLAKSWDISD DGVVYTFHLR DNLVWSDGVS
ITAAE

t6-21.aa

CVNESNRNKLVFKLNIGSEPATLDAQLINDTVGSGIVSQMFLGILDGDPRGGYRPLAKSWDISDDGVVYTFHLR
DNLVWSDGVSITAAE

f6-27.nt

TAAAGAAAAG CTTGCATAAA AAGTATAACA AATTCTTTAA TAATTAAAAT CAAAAAGAAT
ATAATTATTG CACTAAAATT AAATTATAC AGTTATATAG AATCACTTAA GGAACAAAAA
ATGAAATACC TTAAAAACAT TTCCCTTATT TTGTTAATT TAGGTTGCAA ATCCATCCCA
AATGGTAATT TCAATCTACA CGATACAAAC CATAAATTAG GAAAACCTAA ATTTCAGAA
GACTCGATAA TAAGCAGAAA TTATGATAAT AAAATATCCA TTGTTGGAGT ATACAACCT
TTAACAGAAA AAGAAAATT TAAAGTCAT ATTTCATCA AAAAAAAAGG ATTACAAATA
GATCCTGAAA ATATTGAT AAATGAAGAA AAAATTAATT ATTCAAAATA TAAAGCAGAA
CTCAAAGTAA AATCTAGCTT TAATAAAAGC ATTATCAGTA TTTCACTAAC TAATTCAAGA
GATCTATTAA CCTACATTAA CGATAAAAGC ACAGGGAAAT ACATTAACAT TGACTTTAAG
GACAATTGGA ACGTATCGCA CAGTATAAAA TTAAATAAGG AGTATATTAGT AGCATATATA
ACAGATTTG ATAAAGAAAT TAAATATCT AAAATATTT TGCAAAAACG TATTGATAAT
AGAAAAATTG AAATTGAAAA AAAGAGCTT AAAACAGAAAT ATAATGAAAT AGAGGATTAT
TACATCTACA GTATGAAAAT TCCAAAATTA TTGAAAAAT CAGACGCTCC CTCTGAAACT
TACGAAAACAT TTGTTATAGC AAATTATTAC CCCGTGAAA ATTAAATAT ACTGTTTG
AATTAAAGCT TATACTCTGA TAAATTACGC TTTCTAAACT CTATTATGA TGAGAATGAT
AGAAAATTAA AAATGGAGCC TCCGTGAGA GCCTTAAAGA ATTCAAAAC AATAAAAGAA
ACATTAAATA TAGTATTAAAG TCCCTAAAAA ATAATAGAGC TAGCAAAAAA CATTGAAAAA
GATATTACTC TAAAATTAAA ATCTTACGGA GAAAAGGGAG AATTACACATT TGAAATATAT
AAACCACTTC TTTTAAAATT CTTAAAAGAA GTAGATCATT GCATAAAAAA TTTGCAATCA
AGTAGGCATA AATTAA

t6-27.nt

TTGCAAATCCATCCAAATGGTAATTCAATCTACACGATACAAACCATAAATTAGGAAAATCAAGAA
GACTCGATAATAAGCAGAAATTATGATAATAAAATATCCATTGTGGAGTATACAACCCCTTAACAGAAAAAGAAA
ATTTAAAGTCATATTTCATCAAAAAAAAGGATACAAATAGATCCTGAAAATATTGATAATGAAGAAAA
AATTAAATTATTCAAAATATAAGCAGAACTCAAAGTAAATCTAGCTTAAATAAAGCATTATCAGTATTCACTA
ACTAATTCAAGAGATCTATTAAACCTACATTACGATAAAAGCACAGGGAAATACATTAACATTGACTTTAAGGACA
ATGGAACGTATCGCACAGTATAAAATTAAAGGAGTATATTAGCATATAACAGATTGATAAGAAAAT
TAAAATATCTAAAATATTGCAAAAACGTATTGATAATAGAAAATTGAAATTGAAAAACAGAGCTTAAACAA
GAATATAATGAAATAGAGGATTATTACATCTACAGTATGAAAATTCCAAAATTATTGAAAATCAGACGCTCCCT
CTGAAAACCTACGAAACATTGTTATAGCAAATTATTACCCCTGTGAAAATTAAATATACTGTTTGAATTAAAG
CTTATACTCTGATAAAATTACGCTTCTAAACTCTATTATGATGAGAATGATGAGAAAATTAAATGGAGCCTCCT

TABLE 1. Nucleotide and Amino Acid Sequences

GTGAGAGCCTAAAGAATTCAAAAACAATAAAAGAACATTAATATAGTATTAAGTCCTCAAAAAATAATAGAGC
TAGCAAAAACATTGAAAAGATATTACTCTAAAATTTACGGAGAAAGGGAGAATTACACATTGAAAT
ATATAAACCACTTCTTTAAAATTCTAAAAGAAGTAGATCATTGCATAAAAATTGCAATCAAGTAGGCATAAA
TTT

f6-27.aa

RKACIKSITN SLIIKIKKNI IIALKLNLYS YIESLKEQKM KYLKNISLFL LILGCKSIPN
GNFNLHDTNH KLGKLKFQED SIISRNVDNK ISIVGVYNPL TEKENFKVNI FIKKKGLQID
PENILINEEK INYSKYKAEL KVKSFKNSI ISISLTSRD LLTYIYDKST GKYINIDFKD
NWNVSHSIKF NKEYILAYIT DFDKEIKISK NILQKRIDNR KIEIEKTELK TEYNEIEDYY
IYSMKIPKLF EKSDAPSETY ETFVIANYYP CENLNILFLN LSLYSDKLRF LNSIYDENDR
KLKMEPPVRA LKNSKTIKET LNIVLSPQKI IELAKNIEKD ITLKLKSYGE KGEFTFEIYK
PLLLKFLKEV DHCINKLQSS RHKF

t6-27.aa

CKSIPNGNFNLHDTNHKLGKLKFQEDSIISRNVDNKISIVGVYNPLTEKENFKVNI FIKKKGLQIDPENILINEEK
INYSKYKAELKVKSFKNSIISISLTSRDLLTYIYDKSTGKYINIDFKDNWNVSHSIKFNKEYILAYITDFDKEI
KISKNILQKRIDNRKIEIEKTELKTEYNEIEDYYIYSMKIPKLF EKSDAPSETY ETFVIANYYPCENLNILFLNLS
LYSDKLRFLNSIYDENDRKLKMEPPVRAKNSKTIKETLNIVLSPQKIIIELAKNIEKDITLKLKSYGEKGEFTFEI
YKPLLLKFLKEVDHCKINKLQSSRHKF

f6-5.nt

TAAATGAAGA AGTTTTAAT ATCCGTTAT TTTTATTGT TTTATGGTG TTCAACTATA
TCTTTGGTAA AAATACCGA AAAAGATAAA ATAATTTAA CTGTTTATC ATCTTTAATG
AATTATCCTG ATTTGAAGAT TTCAAATTT AAAATAAAAG ACTACGAACA TTTGCATTAT
TCATCTGATT TTGAAAGCTT GAGTGATACT AAAATAGTG CTTATATTAA CGTTGATGAA
TCTAGTTCA ATAATAATAT TAATTTATT AAAGATCTT TTATTATAA TAAGAAATTA
TATAGAATAC TTATTGCTTA TAGCTTGACC CAAGGTGCAT CTTTTAAGGC AGAAGTTTA
TCTTATCTTG AAAAACAAAA AATTATGAAA AATTTTCAT TGAAAATAAA TTTCCAAC
GCTAAAAAAT TTATGGATAA TAAGTATTGG ATTGTAATTG CAAAAAACCA TTTAGATTCT
CTTGTAAAGA GTAAAAATTA TTTAGTCTTG GCGAATGTAA AGATGGAATA TATACTCAA
AAGTTTTAA CTTGA

t6-5.nt

TTGTTCAACTATATCTTGGTAAAATACCAGAAAAGATAAAATTAACCTGTTTATCATCTTAAATGAAT
TATCCTGATTGAGATTCAAAATTAAAATAAAAGACTACGAACATTGCATTATTCATCTGATTGAAAGCT
TGAGTGATACTAAAATAGTGCCTATATTACGTTGATGAATCTAGTTCAATAATAATTAAATTAAAGA
TCTTTTATTATAATAAGAAATTATAGAATAACTTATTGCTTATAGCTTGACCCAAGGTGCATCTTTAAGGCC
GAAGTTTATCTTATCTGAAAACAAAAATTATGAAAATTTTCAATTGAAAATAAAATTCCAAC
AATTATGGATAATAAGTATTGGATTGTAATTGCAAAAACCATTAGATTCTTGTAAAGAGTAAAAT

f6-5.aa

MKKFLISVYF LLFYGCSTIS LVKIKEKDKI NLTVLSSLMN YPDLKISNFK IKDYEHLHYS
SDFESLSDTK NSAYIYVDES SFNNNINFIK DLFYIYKLY RILIAYSLTQ GASFKAEVLS
YLEKQKIMKN FSLKINFPTA KKFDNQYWI VIAKNHLDL VKSKNYLVLA NVKMEYILKK
FLT

t6-5.aa

CSTISLVKIKEKDKINLTVLSSLMNYPDLKISNFKIKDYEHLHYSSDFESLSDTKNSAYIYVDESSFNNNINFIK
LFIYKLYRILIAYSLTQGASFKAEVLSYLEKQKIMKNFSLKINFPTAKKFMDNQYWIVIAKNHLDLSLVSKN

TABLE 1. Nucleotide and Amino Acid Sequences

f7-30.nt

TAGAGACGAA GTCACAAGCA AAATGTTAA AGATTTACAA AATCAAGTTC AAGGGGGCAA
 ATAATGAAAA ATTTAAAGAC AAAAATTAAT TTTTAGGGA TATTTGGCT ACTGTTACTA
 TTTCTTCTT GCGAATCAAT ACCATCACTT CCCCAAAAC CAACCTAAC AAACAAAGAA
 GATATTGAA ATTTAATGCT CGATGAAGCA GAACTTTTA GATACTAAC CGCACTAAAT
 GTTGGCTT TGACTGTAA ATCTTATGTG ATCAAATACT ATCCTAATGA CAAATTCCT
 GTGTTGAA ATTTGATCC CGTGTGCG GATGAAAATG GAACTAAAGA AACAAATATA
 CTAAGAACATC GAATTACCTA CTACAATCGA TACATAGAA AAACCGAAC GATTGTATT
 GGGTGTACAA AAAATACAG CAGAAGATAA

t7-30.nt

TTGCGAATCAATACCATCACTTCCCCAAAAACCAACCCCTAACAAACAAAGAAGATATTGAAAATTAATGCTCGAT
 GAAGCAGAACTTTTAGATACTCAACCGCACTAAATGTTGGCTTGTGACTGAAATCTTATGTGATCAAATACT
 ATCCTAATGACAAATTCTGTGTTGAAAATTTGATCCCGTGTGCGATGAAATGAACTAAAGAAACAAA
 TAACTAAAAATCGAATTACCTACTACAATCGATACATAGAAAAACCGAACCGATTGTATTGGTGTACAAA
 AAATACAGCAGAAGA

f7-30.aa

RRSHKQNVKR FTKSSSRGQI MKNLTKINF LGIFWLLLLF LSCEISPLP QKPTLTNKED
 IENMLMDEAE LFRYSTALNV WLLTVKSYVI KYYPNDKFPV FENFDPVFGD ENGTKETNIL
 KNRITYYNRY IEKTEPIVFG CYKKYSRR

t7-30.aa

CESISPLPQKPTLTNKEDIEMLMDEAELFRYSTALNVWLLTVKSYVIKYYPNDKFPFENFDPVFGDENGTKETN
 ILKNRITYYNRYIEKTEPIVFGCYKKYSRR

f76-1.nt

TGAATATTAA TAATAAAAAA AGGAGTAACA ATGAAAATTA TCAACATATT ATTTGTTTG
 TTTTACTAA TGCTAACCGG CTGTAATTCT AATGATACAA ATACCAAGCA GACAAAAAGC
 AGACAAAAGC GTGATTTAAC CCAAAAGAA GCAACACAAG AAAACCTAA ATCTAAATCT
 AAAGAAGACC TGCTTAGAGA AAAGCTATCT GATGATCAAA AAACACAACT TGACTGGTTA
 AAAACCGCTT TAACTGGTGT TGGAAAATTT GATAAATTCT TAGAAAATGA TGAAGGCAA
 ATAAATCAG CACTGAACA TATAAAGACT GAACTTGATA AATGTAATGG AAATGATGAA
 GGAAAAAACCA CCTTCAAAAC TACCGTTCAA GGGTTTTA GCGCGGCAA TATAGATAAT
 TTGAGATC AAGCAACTGC TACCTGCAAT TAA

t76-1.nt

CTGTAATTCTAATGATACAAATACCAAGCAGACAAAAAGCAGACAAAGCGTGATTTAACCAAAAGCAACA
 CAAGAAAAACCTAAATCTAAATCTAAAGAAGACCTGCTTAGAGAAAAGCTATCTGATGATCAAAACACAACCTG
 ACTGGTTAAAACCGCTTAACTGGTGTGAAAATTTGATAAATTCTTAGAAAATGATGAAAGGCAAATTAATC
 AGCACTTGAACATATAAAGACTGAACTTGATAAATGTAATGAAATGATGAAAGGAAAACACCTTCAAAACTACC
 GTCAAGGGTTTTAGCGGGGCAATATAGATAATTTGAGATCAAGCAACTGCTACCTGCAAT

f76-1.aa

ILIIKKVTM KIINILFCLF LLMLNGCNSN DTNTKQTKSR QKRDLTQKEA TQEKPKSNS
 EDLLREKLSD DQKTQLDWLK TALTGVGKFD KFLENDEKSI KSALEHIKTE LDKCNGNDEG
 KNTFKTTVQG FFSGGNIDNF ADQATATCN

t76-1.aa

TABLE 1. Nucleotide and Amino Acid Sequences

CNSNDTNTKQTKSRQKRDLTQKEATQEKP KSKSKEDLLREKLSDDQKTQLDWLKTALTGVGKFDKFL ENDEGKIKS
ALEHIKTELDKCN GNGNDEGKNTFKTTVQGFSGGNIDNFADQATATCN

f8-10.nt

TAAGTAAGGA GAATATTTAT GAAATATAAT ACGATTATAA GCATATTGT TTGTTGTT
TTAAGTGC TT GCAATCCAGA TTTTAACACA AATAAGAAA GAACTCTAAG TAAGGGATA
ATTTCAAATC AAGATGCAGA TTCTGATAAA ATAATAAAA ATAATTACT TGATGATT
ATAAATTAA TAGAAAAAGC GAATGCAGAT AGAGAAAAT ATGTA AAAAAT AATGGAAGAA
GAACCTTCGG ATCAATATGG AATGTTGGCT GTTTTGAG GTATGTATTG GGCAGAATCA
CCACGGAAAT TAATATCTGA TACAGGTAGT GAGAGATCTA TTAGGTATAG AAGGCGTGT
TATAGTATT TATTAAATGC TATTGAAACT AATGAATTAA AGAAATTTC AGAAATTAGA
ATACTGTCAA TAAAAGTACT AGAAATATT AGCCTATTG ATCTATTG AAGTACTCTT
GATGATGTGG TTGTTCACTT ATATTCCAAA AAAGATACTC TAGGTAACAG ATGATATTCA
AATTTAAAAA GACTAAAAA TTTGTTGAA AAATTATTAT CTATAAAAAC AATGTTCA
AAGATGTCAA AACGCTTTT ATTGGATTAT CAAAATAATG AAAATTTC AAAAAACAGAT
AACGCCAAGC TTGGATCTTA TGTGGTTGCA CTTCCAATC AAATTCAAGA AAAATATAAT
GAAGCAGAAA GGCTGAAAAG CGAGATAATT TTAATATATA CCCTTAA

t8-10.nt

TTGCAATCCAGATTTAACACAAATAAGAAAAGAACTCTAAGTAAGGGATAATTCAAATCAAGATGCAGATTCT
GATAAAATAATAAAAATAATTACTTGATGATTTAATAAATTAAATAGAAAAGCGAATGCAGATAGAGAAAAT
ATGTAAAAAAATGGAAGAAGAACCTCGATCAATATGGAATGTTGGCTGTTGGAGGTATGTATTGGCAGA
ATCACCACGGAAATTAATATCTGATACAGGTAGTGAGAGATCTATTAGGTATAGAAGGCGTGT
TTAAATGCTATTGAAACTAATGAATTAAAGAAATTTCAGAAATTAGAATACTGTCAATAAAAGTACTAGAAATAT
TTAGCCTATTAAATCTATTGGAAGTACTCTGATGATGTGGTTGTTCACTTATATTCAA
TAAACTAGATATTCAAATTAAAAGACTTAAAATTGTTGAAAATTATTATCTATAAAAACAATCGTTCA
AAGATGTCAAACGCTTTTATTGGATTATCAAATAATGAAAATTTATAAAAACAGATAACGCCAAGCTGGAT
CTTATGTGGTTGCACTTCCAATCAAATTCAAGAAAATATAATGAAGCAGAAAGGCTGAAA

f8-10.aa

VRRIFMKYNT IISIFVCLFL TACNPDFNTN KKRTLSKGII SNQDADSDKI IKNKLDDLI
NLIEKANADR EKYVKKMEEE PSDQYGMALV FGGMYWAESP RELISDTGSE RSIRYRRRVY
SILLNAIETN ELKKFSEIRI LSIKVLEIFS LFNLFGSTLD DVVVLHLYSKK DTLGKLDISN
LKRKLNLFEK LLSIKTIVSK MSKRLLLDYQ NNENFIKTDN AKLGSYVVAL SNQIQEKYNE
AERLKSEIIL IYTL

t8-10.aa

CNPDFNTNKKRTLSKGII SNQDADSDKI IKNKLDDLI NLIEKANADREKYVKKMEEPSDQYGMALVFGGMYWAE
SPRELISDTGSE SRSIRYRRVY SILLNAIETN ELKKFSEIRI LSIKVLEIFS LFNLFGSTLD DVVVLHLYSKD LG
KLDISNLKRLKNLFEK LLSIKTIVSKMSKRLLLDYQNNENFIKTDNAKLG SYVVAL SNQIQEKYNE AERLK

f8-14.nt

TAATATATAT TCTTGATTA GGGAAAGGAG AGTATTTTA TGAAAAAAA AATGTTTAA
TATACATTGT TAACGATAGG ATTGATGTCT TGTAATCTAA ATTCTAAATT ATCTGGTAAT
AAAGAGGAAC AAAAATAA CAATGATATA AAAGAAGCTT TAAATGGCGT TCAAGAAAAT
GCTATTAAATA ATTATATGG AAATAAAA GAAAAGGAG ATTATGTTAA AAATTGGAA
AAATTGAAAG ACAAGGGTTT AGACGTGACC ACCCTCCCT TAGAACCTGT AGTGGCGCCC
TCCGTAGAAT CTGCGGTGTC TTTAGGAGAA TCTAATAATA GGATTGGTAT ACCAACCA
TCAATTGAGC ATAATCAA AAAAGAGATA AAAGAAGAGG ATTGTTTCCC TTCTACTGAG
GAAGAAAAGC AAGCGGATAA AGCAATTAA GATATAGAGA ATCTTATTG AGAATCTGGA

TABLE 1. Nucleotide and Amino Acid Sequences

TTTCCCCAGT TAATTGAGAA TGTGTGCTCA CTTAACATG AATATACTTT AATAAGAAGT
 GATTTTTATG ATGTGATAAC TAAGATTCAAG AATAAAAAAA TATCACTAAT GAAAAATTCT
 CATAATAATA GAAATAAAAT AAGGGAACTA GTACAATTGC AAAATAATT AAAGATAGGA
 GACGAACCTG ATAAAATTAT CGGTTGCATT GATACTGCAG ACAAGAGAT AAGATCTGCC
 GCTTTCTTT TTGATGAAGC TAAGGAAAGC TTAAAAGAAG GTATTATTAA AAGATTGGAA
 AAAAGTAAAA ATAGGGCAGC ATCACAATT A TCTAAAAGG CTTAAATAG AGCAGAGGAT
 GCTTTAAGGT GCTTAGAAAA TTATTCTCT AAAAGGAGTGTG AGGCAATAGG AAGAAGAAGC
 TTTATAAAAG AAGTTGTTGA ACAGGCAAAA AATGCTTTAA GTAAAGTCTTA A

t8-14.nt

TTGTAATCTAAATTCTAAATTATCGGTAAATAAGAGGAACAAAAAAATAACAATGATATAAAAGAAGCTTTAAAT
 GCGTCAAGAAAATGCTATTAATAATTATGAAATAAAAAAGAAAAAAAGATTTATTAAAAATTCCGAAA
 AATTGAAAGACAAGGGTTAGACGTGACCACCCCTCCCTAGAACCTGTAGTGGCGCCCTCCGTAGAATCTCGG
 GCTTTAGGAGAATCTAATAATAGGATTGGTATACCAACCATTCAATTGAGCATAATCAAAAAAAAGAGATAAAA
 GAAGAGGATTTTCCCTACTGAGGAAGAAAGCAAGCGGATAAGCAATTAAAGATATAGAGAATCTTATTG
 GAGAATCTGGATTCCGAGTTAATTGAGAATGTGCTCACTAAACATGAATATACTTTAATAAGAAGTGT
 TTATGATGTGATAACTAAGATTCAAAATAATCAACTAATGAAAATTCTCATAATAATAGAAATAAAAATA
 AGGGAACTAGTACAATTGCAAAATAATTAAAGATAGGAGACGAACCTGATAAAATTATGGGTTGCATTGATACTG
 CAGAACAAAGAGATAAGATCTGCCGTTCTTTGATGAAGCTAAGGAAAGCTTAAAGAAGGTATTATAAAAG
 ATTGGAAAAAGTAAAATAGGCAGCATCACATTCTAAAGGCTTAAATAGAGCAGAGGATGCTTAAGG
 TGCTTAGAAAATTATTCTTCTAAAAAGGTGAGGAATAGGAAGAAGCTTATAAAAGAAGTTGTTAACAGG
 CAAAAAAATGCTTTAAGTAAGTCT

f8-14.aa

YIFLIKGES IFMKKKMFY TLLTIGLMSC NLNSKLSGNK EEQKNNNNDIK EALNGVQENA
 INNLYGNKKE KKDFIKNSEK LDKGLDVTT LPLEPVVAPS VESAVSLGES NNRIGIPTIS
 IEHNQKKEIK EEDFPSTEE EKQADKAIKD IENLIGESGF PELIENVCSL KHEYTLIRSD
 FYDVITKIQN KKISLMKN SHNNRNRKIRELV QLQNNLKIGD ELDKIMGCID TAEQEIRSA
 FFFDEAKESL KEGIIKRLK SKNRAASQLS KKALNRAEDA LRCLENYSSK KGEAIGRRSF
 IKEVVEQAKN ALSKS

t8-14.aa

CNLNSKLSGNKEEKNNDIKEALNGVQENA INNLYGNKKEKKDFIKNSEKLKDGLDVTLPLPVVAPS
 VESAVSLGESNNRIGIPTISIEHNQKKEIK EEDFPSTEEEKQADKAIKD
 IENLIGESGFPELIENVCSL KHEYTLIRSD
 YDVITKIQNKKISLMKN SHNNRNRKIRELV QLQNNLKIGD ELDKIMGCID TAEQEIRSA
 FFFDEAKESL KEGIIKRLK SKNRAASQLSK KALNRAEDA LRCLENYSSK
 KGEAIGRRSF
 IKEVVEQAKN ALSKS

f01A.nt BB001

TGATTAATTTTTAAGGATTACGTTGAAAAGAAACAAAATTGGAAAAGCTTAAACTGTTCAAATAACTT
 TACTGTTCTCATGCTCTTTATTCTAAATCAAACACAGAAGCGATAAGTGAATTACAATCAAGCCCTATTAA
 ACTTGGAAAAATTAAAGTTTACAAAAACAGAAAAGATTGTAAGCACC AAAACTCTCAAACAAAGC
 CAGTTCTTAAAAATGAAAAGAAAATAATTAAAAATTGACACAAGAATTGATGAGAATGAAAATTGATTA
 ATAAAATAGGTCAAATATGAAATGTTGCTCAAACAATAAACACGGATATTCAA
 AAAATCGAACCTAATGATCA
 ATTGGAATAAAACTTATTACAGAAAAAGACAATAATTGACTTATGTTAAAGACAATCGACTT
 AGAAGATTATTACTCATCTTAAATTGATGAAAATAAAATTGCCACAATACTCGCGAAACAT
 CAAGCTCAAACGACTACCATTACACACTTATTGGTTAATTGGACAGGATTAAATCCAAGAAGCATTGA
 AAGCGCTGTAATATTAACTAAAGACGAGCAAAGCGCTAATTAA
 ATTGAAACAAAAACAGTAAAGAG
 ATTCAAGGAAAATTGAAAAGCTAATGCAAGAGAGAATTGATGAAAATTGTCGATAACATTATTGGCGAAT
 ATGACAAAATACGGGAGGATGCAAAGCTGATGAAAATTCTGGAGAAGTAATAAGGTTGGATACGAGCATTGA
 ACTCGACTCAAATAAAAGTATGCAAATTAAACAATATTGAAACACCGCTAAAACCTGTTGACCACATACAC
 TACTAA

TABLE 1. Nucleotide and Amino Acid Sequences

t01A.nt BB001

TGCTCTTTTATTCTAAACAAACACAGAGCATAAGTGAATTACAATCAAGCCCTATTAACCTGGAAAAAA
 TAAAGTTTACAACAGAAAAGATTGTAAGCACCACAAACTTCAAAACTACAACAAAGCCAGTTCTTAA
 AAATGAAAAAGAAAAATAATTAAAAAAATGCACAAGAATTGATGAGAATGAAAATTGATTAATAAAATAGGT
 CCAAATATCGAAATGTTGCTAAACAATAAACACGGATATTCAAAATCGAACCTAATGATCAATTGGAATAA
 ATAAAACCTTATTCAACAGAAAAAGACAATAATTGACTTATGTTAAAGACAATCGACTTAGAAGATTATT
 TTACTCATCTTAAATTATGATGAAAATAAAATCAAAAAATTAGCCACAATACTCGCGAACATCAAGCTCAAAC
 GACTACCATTACACACTATTGGTTAATTGGACAGGATTAAAATCCAAGAACGATTGAAAGCGCTGTTA
 ATATTAACTAAAGACGAGCAAAGCGCTAATTTAATTGAGAACAAAACAGTAAAGAGATTCAAGGAAA
 TTTGAAAAACTAATGCAAGAGAGAAATTGATGGATAAAAATCGTCATAACATTATTGGCAATATGACAAAAAAT
 ACGGGAGGATGCAAAGCTGATGAAAAATTCTCGGAGAAGTAATAAGGGTTGGATACGAGCATGAACTCGACTCAA
 ATAAAAGTATGCAAATTAAACAATTGAAACACCGCTAAAACCTGTTGACCACATACACTAC

f01A.aa BB001

LIFFKDYVLKRNIWKTALKFQITLLFSCSFYSKSNNTAISELQSSPIKLGKIKVLQKTEKIVSTQNLQNLQQSQ
 FFKNEKEKIIKKIAQEFDENEKLINKIGPNIEMFAQTINTDIQKIEPNDQFGINKTLFTEKKDNNIDFMLKDNRLR
 RLFYSSLNYDENKIKKLATILAQTSSSNDYHYTLIGLIFWTGFKIQEAFESAVNILTKEQKRLIFNFRKTKEI
 QENFEKLMQERNNSWIKIVDNIIGEYDKNTGGCKADGKILGEVIRVGYEHELDNSKSMQILNNIETPLKTCCDHIIHY

t01A.aa BB001

CSFYSKSNNTAISELQSSPIKLGKIKVLQKTEKIVSTQNLQNLQQSQFFKNEKEKIIKKIAQEFDENEKLINKIG
 PNIEFAQTINTDIQKIEPNDQFGINKTLFTEKKDNNIDFMLKDNRLRLFYSSLNYDENKIKKLATILAQTSSSN
 DYHYTLIGLIFWTGFKIQEAFESAVNILTKEQKRLIFNFRKTKEI QENFEKLMQERNNSWIKIVDNIIGEYDKN
 TGGCKADGKILGEVIRVGYEHELDNSKSMQILNNIETPLKTCCDHIIHY

f02A.nt BB002

TAATTAATACTGGTTAATTATAAGGAGAGTATTGAAAAAAAGCAAACAAATATAATCAAGATTAATATTA
 TTACAATGATATTAACTTAATTGATCTCATGTGACCTTTAACAAAATCAATCCAAGGCAAATGAAAACAC
 CAAGCTAAAAAAACACCAGACTGAAAAACCCGCAATCCAGGGAAACATCCAATTTAAAGATAAAATCT
 GGAGACCTTGGCGCTTCTGATGAAAATTATGGAACTACCGCTTCAGAGCTAAAGCAATTGTAAGGAGCTAG
 AAGATGAAAAAAATCAATACGATAACAAATAGCAAATTACTAATGAAGAATCTAACCTATTAGATACTTATAT
 TCGGGCTTATGAACTAGCTAACGAAAATGCTTTAAAAGATTCTTCTTCAATCTTCTTCAATTAGATTATAAA
 AAAGAAAACATAGAGACATTAAAGAAATTCTGAAAAACTCATAAAATTACGAAAACGACCCAAATTGCTG
 CAAATTCCCTTATGCATAGCGCTGGATATTCAATTAAAACCTGGAAAAGCACTTAAATCAATAATGAAAAC
 GGACACTCTAACGAAAAGAAATTCAAAAGAAGATTAGAGGCGTTGCTAGAACAGTAAAATCTGCCTTACAGCTA
 CAAGAAAAGTTAAAAAACCTAAACAAAACCTTGAAGATTACCGTAAAATACTAACACATTCAAGAAAATA
 AAGTACTAGCAGAACACTTAAATAATATTACAAAGACTCTGATTCTTACAATCTGCCTTTATTAA

t02A.nt BB002

TGTGCACCTTTAACAAATCAATCCAAGGCAAATGAAAACACCAAGCTTAAAAAAACACCAAGACTGAAAAAC
 CCGCCAATCAGGGAAAACATCCAAAATTAAAGATAAAATCTGGAGACCTTGGCGCTTCTGATGAAAATTAT
 GGGAACTACCGCTTCAGAGCTAAAGCAATTGCTAGAGCTAGAACATCGATATAACAAATA
 GCCAAAATTACTAATGAAGAATCTAACCTATTAGATACTTATTCGGGCTTATGAACTAGCTAACGAAAATGAAA
 AAATGCTTTAAAAAGATTCTTCTTCACTTTAGATTATAAAAAGAAAACATAGAGACATTAAAGAAATTCT
 TGAAAACCTCATAATAATTACGAAAACGACCCAAAATTGCTGCAAATTCTTCTTATGCATAGCGCTGGATATT
 CAATTAAAACGGAAAAGCACTTAAATCAATAATGAAAACCTGGACACTCTAACGCAAAGAAAATTCAAAAGAAG
 ATTAGAGGCGTTGCTAGAACAGTAAAATCTGCCTTACAGCTAACGAAAAGTTAAAAAACCTAAACAAAAC
 TCTTGAAGATTACCGTAAAATACTAACACATTCAAGAAAATAAGTACTAGCAGAACACTTTAATAAAATTAC
 AAAGACTCTGATTCTTACAATCTGCCTTTAT

f02A.aa BB002

TABLE 1. Nucleotide and Amino Acid Sequences

LILVLIYKESILKKAKLNIKINIITMILTLICISCPFNKINPKANENTKLKKNTRLKKPANPGENIQNFKDKSG
 DLGASDEKFMGTTASELKAIGKELEDRKNQYDIQIAKITNEESNLDTYIRAYELANENEKMLLKRFLLSSLDYKK
 ENIETLKEILEKLINNYENDPKIAANFLYRIALDIQLKLEKHLKSINEKLDLTSKENSKEDEALLEQVKSLQLO
 EFKKTLNKTLEDYRKNTNNIQENKVLAEHFNKYYKDSDSLQSAFY

t02A.aa BB002

CAPFNKINPKANENTKLKKNTRLKKPANPGENIQNFKDKSGDLGASDEKFMGTTASELKAIGKELEDRKNQYDIQI
 AKITNEESNLDTYIRAYELANENEKMLLKRFLLSSLDYKKENIETLKEILEKLINNYENDPKIAANFLYRIALDI
 QLKLEKHLKSINEKLDLTSKENSKEDEALLEQVKSLQLOEKFKKTLNKTLEDYRKNTNNIQENKVLAEHFNKYY
 KDSDLQSAFY

f03A.nt BB006

TGATTTAATGTAATTAACTACCGCTAAAAAAGGCTTAAATGGTATAAAGGAAGAGCTAATGGTATTTA
 GAACATATAAACATTGGAACATAATAATGCTGCCATGTTAATGCTGAGTTGCGCTTTTTAAGAAACCACAATC
 TGACATCAAGACAGCAATACTGGCAACCAATAAGCGATGAAAAATTACATTAAATATCAGGAAAATTCAAAT
 AAAAATTGCCAATCATAAATAGTAATCATGACGTAACCTGGATAAAAACAAAGGCAATGACAATCTTAGGCGAAG
 ATGGAAAAGAAATACCGAATTAAAAACAAATTGGATATTCTTATATAATATCTCTGTAAAAATGGATGGAAA
 ATATAGTTATTACGGCTCATTATAATACCTTTGAAACAACTAAAAATGGAGATGATGAATATGAAATTGAAGAT
 GTTAAATTGTAACAGCTGGTCCACCCCTAGAACTTAAAATCTCTTAGCTGTTGAAAATTCAAAGAAGAAG
 GATATGTTACTGCATACCCATTGGAATTGATGAGTGACGAGATTAAAATGCTTTAAATTAAACATATAAAA
 TGGTCAATTGGAATTATATGCTGCAGATTAACTGTCAAAATAACTACTCAAGAAAACCTAAATTATAAAAATT
 TCTCTTAATTCAAATTATTGAATTTTAAAAGAAGTGTAAAAGAAAATTCTATATTAAAAGACATAGCTG
 GAGATTATTGAAGATATAA

t03A.nt BB006

TGCGCTTTTTAAGAAACCACAATCTGTACATCAAGACAGCAATACTGGCAAACCAATAAGCGATGAAAAATTAC
 ATTTAATATCAGGCAAATTCAAATAAAAATTGCCAATCATAAATAGTAATCATGACGTAACCTGGATAAAAAC
 AAAGGCAATGACAATCTAGGCGAAGATGAAAAGAAATACCGAATTAAAACAAATTGGATATTCTTATATA
 ATATCTCTGTAAAATGGATGGAAAATATAGTTATTACGGCTCATTATAATACCTTTGAAACAACTAAAATG
 GAGATGATGAATATGAAAGATGTTAATTGTAACAGCTGGTCCACCCCTAGAACTTAAAATTCTCTTT
 AGCTGTTGAAAATTCAAAGAAGGATATGTTACTGCATACCCATTGGAATTGATGAGTGACGAGATTAAA
 AATGCTTTAAATTAAACATATAAAATGGTCAATTGGAATTATGCTTGAGATTAACTGTCAAAAATAACTTA
 CTCAAGAAAACCTAAATTATAAAATTCTCTTAATTCAAATTAAATTATTGAATTTTAAAAGAAGTGTAAAAGA
 AAATTCTATATTAAAAGACATAGCTGGAGATTATTGAAGATATA

f03A.aa BB006

FNVNFNYRLKKALNGIKEEDLMVFRYKHELEIMLPMMLSCAFFKKPQSVHQDSNTGKPISEKHLISGKISNK
 KLPIINSNHDVTWIKTKAMTILGEDGKEIPEFKNKGYSYIISPVKMDGKYSYYASLLILFETTKNGDDEYEIEDV
 KFVTAGSTLELKNSLLAVENSQEEGYVTAYPFGILMSDEIKNAFKLTYKNGHWNYMLADLTVKNKLTQETKLYKIS
 LNSKLIIEFLKEVLKENSILKDIAGDLFEDI

t03A.aa BB006

CAFFKKPQSVHQDSNTGKPISEKHLISGKISNKKLPIINSNHDVTWIKTKAMTILGEDGKEIPEFKNKGYSYI
 ISPVKMDGKYSYYASLLILFETTKNGDDEYEIEDVFKVTAGSTLELKNSLLAVENSQEEGYVTAYPFGILMSDEIK
 NAFKLTYKNGHWNYMLADLTVKNKLTQETKLYKISLNSKLIIEFLKEVLKENSILKDIAGDLFEDI

f04A.nt BB011

TAATTACCAAAGATAAGTAAACTTGCACAAATAAAACTACACGTATTGAAAGTAGATTGAAATTCCATTATTTA
 TATATAATGGCACTAAATATCTGAAAATGAAGGAGAAGCGGGTGGGCAATAAAATTCTTATATTTCAGTGGTTT
 AATTAAATAGTTGGTTGCGACTGGGAACATTAAAGATAAAAGTACAGAAATTCCAAGCTATTAAAGAACGGAC

TABLE 1. Nucleotide and Amino Acid Sequences

AAAGATAAGACTAAAATCAAGATAGAATAGAATTGGGTGAAGATAATTTGTATCTAAAATAATATGTCTACTA
 CTGATACGGCATTACTAGTTAGGAAGTCTAACAACTTGGATTTAATTAAATCGTCACAGCGGGTCAGTGAACC
 ACTTATAATCTCAAATGAGAAAGCCATAGCTACTCAAGCAAAGTAGATTAAATGAACAACATTAATGTTACTATA
 ATAAACCCAAAACCAGCTCAAATTTGGAAATTCTTAAACAATACTACTACTGAAGATAGTGTGAAGTTTTAT
 CAATTGAAAACCAAGAGTGGCTTATTAGTAAAAGATTGCCCAGTAAGTTGGAAAATTAGAAAGCTTCTAAA
 AACACAACACGAAAAGAAGCTTTAAGACGGCTAAAACATACAAAGTCTCATTAGTAATTCCAATATGGTAAA
 GAAATTATTAAGTTAAGGAAGAATTACAAACTTTATAATTGTTGAAGGCATACAACAAAATTCCATAGTC
 AAAGGAATTCTTATAAAAGATACTAAATTGGGAAAATAGACAAAAAATGCAGTTATTTAAATCCTTTTC
 ATCTATAGAGAAAGAATTAGAGATTGAAATTATAAGTTGNGTGAATCCTAAAGTAATTTCAAATTGCAGATGTT
 AGCTGGAATAATGCAAACCTCTTTAAAGAATCTATAGAAAATTAAATCAGGCAATTGAAAAAGGTATGACA
 ATGAGAGTAGAAAGCAAGGTCAAATTGGTGGACCTGCTAATAGATGGATAAAAATCAAGCTGACAATTGCTAA
 GGATGCAAAGTATAAGGCAGAACATTCAAGGAAATTGATTGGAAAATGCAGCCAATTAGATAGTTGTTCA
 AATGAAAAAGAAGCTAAAAGCTATTAGAAGAAATTAAAAAGATTGTAACGAAATTGGTATTAGCCTATAA

t04A.nt BB011

TGCGACTGGGAACTATTAAAGATAAAAGTACAGAAATTCCAAGCTATTAAGAACGGACAAAGATAAGACTAAA
 ATCAAGATAGAATTGTTGAAGATAATTGTTATCTAAAATAATATGTCTACTACTGATACGGCATTAC
 TAGTTAGGAAGTCTAACAACTTGGATTTAATTAAATCGTCACAGCGGGTCAGTGAACCACCTATAATCTCAAAT
 GAGAAAGCCATAGCTACTCAAGCAAAGTAGATTAAATGAACAACATTAATGTTACTATAATAACCCAAAACCAG
 CTCAAATTGGGAAATTCTTAAACAATACTACTACTGAAGATAGTGTGAAGTTTTATCAATTGAAAACCAAGA
 GTGGCTTATTAGTAAAAGATTGCCCAGTAAGTTGGAAAATTAGAAAGCTTCTAAAACACAACACGAAAAA
 GAAGCTTTAAGACGGCTAAAACATACAAAGTCTCATTAGTAATTCCAATATGGTAAAGAAATTATTAAGTTA
 AGGAAGAATTACAAACTTTATAATTGTTGAAGGCATACAACAAAATTCCATAGTCAGGAAATTCTATTAT
 AAAAGATACTAAATTGGGAAAATAGACAAAAAATGCAGTTATTTAAATCCTTTCATCTATAGAGAAAGAA
 ATTAGAGATTGAATTATAAGTTGNGTGAATCCAAAGTAATTTCAAATTGCAAGATGTTAGCTGGAATAATGCAA
 ACTCTCTTAAAGAATCTATAGAAAATTAAATCAGGCAATTGAAAAAGGTATGACAATGAGAGTAGAAAGCA
 AGGTCAAATTGGTGGACCTGCTAATAGATGGATAAAAATCAAGCTGACAATTGCTAAGGATGCAAAGTATAAG
 GCAGAACATCAGCAAATTGAGATTGGAAAATGCAGCCAATTAGATAGTTGTCAAATGAAAAGAAGCTA
 AAAAGCTATTAGAAGAAATTAAAAAGATTGTAACGAAATTGGTATTAGCCTA

f04A.aa BB011

LPKISKLANKTTRIESRFEISIIFIYNGTKYLKMKEKRVGNKIFYISVVLILIVGCDWGTIKDKSTEISKLLRTDK
 DTKNQDRIELGEDNFVSKNMSTTDGTGITSLSLNNLDLINRSQRVSEPIIISNEKAIATQAKVDMNNINVII
 NPKPAQNLGNLSLNNTTEDSVKFLSIENQEWLISKKILPSKLENLESFLKTQHEKEAFKTAKTIQSLISNSNMGKE
 IIKFKEEYYKLYNLFEGIQQKFHSQRNSFIKDTKFGENRQKNAVIFKSFSIEKEIRDLYKLXEIQSNFQIADVS
 WNNANSLLKESIEKLIQAIKRYDNESRKQGQIGGPANRWDKNQADNFAKDAKYKAEHSANDLENAANYFRYCSN
 EKEAKKLLEEIKKRFVRIGISL

t04A.aa BB011

CDWGTIKDKSTEISKLLRTDKTKNQDRIELGEDNFVSKNMSTTDGTGITSLSLNNLDLINRSQRVSEPIIISN
 EKAIATQAKVDMNNINVIIINPKPAQNLGNLSLNNTTEDSVKFLSIENQEWLISKKILPSKLENLESFLKTQHEK
 EAFKTAKTIQSLISNSNMGKEIIKFKEEYYKLYNLFEGIQQKFHSQRNSFIKDTKFGENRQKNAVIFKSFSIEKE
 IRDLNYKLXEIQSNFQIADVSWNNAKSIEKLIQAIKRYDNESRKQGQIGGPANRWDKNQADNFAKDAKYK
 AEHSANDLENAANYFRYCSN EKEAKKLLEEIKKRFVRIGISL

f05A.nt BB009

TAAATAAAATTGTAGGATAAAAATGAAACAAAATACGAAAACATTTTAAAAAGATTAATTAAACCTATTAA
 TATTTTACTACTAGCATGCTCAAGCGAATCCATATTTCACAATTAGGAATCTGCAAAAATAAAACATGAATA
 CAATATTGGCAGTTCAAGTCAAGAGGAAATTCTCTAGTAGGAGAAACTCTCTACATTGCAGCCATGCATTAA
 TTAAAAAAAGAAAACGGCAAGATTGAAAAATTGATTGAGCAATTCTTATGAGTTATAAAACGACATTGAAATA
 TATCTGGAAAACCTATCTTGTAGCGCAAAACAAAGAAGAAGAATTAGAAGTTGCGAGCTAAATGAAAAGATTG
 GACATTAAAATTAAAAACCGCTAAAGCATATAAATTCTTAAATCCGTAGAAGAGATGGCGTAA

TABLE 1. Nucleotide and Amino Acid Sequences

t05A.nt BB009

TGCTCAAGCGAATCCATATTTACAATTAGGAATCTGCAAAAAATAAAACATGAATACAATATTTGGCAGTT
 CAAGTCCAAGAGGAATTCTCTAGTAGGAGAAACTCTACATTCAGCCATGCATTATTTAAAAAGAAAACGG
 CAAGATTGAAAAATTGATTTGAGCAATTCTTATGAGTTATAACGACATTGTAATATCTGGAAAAACCTAT
 CTTTAGCGCAAAACAAAGAAGAATTAGAAGTTGCGAGCTAAATGGAAAAGATTGGACATTAAAATTTAAAA
 AACCGCTAAAGCATATAAATTCTTAAACCGTAGAAGAGATGGCG

f05A.aa BB009

INCRIKMKQKYENYFKKRLILNLIFLLLACSSSIFSSQLGNLQKIKHEYNILGSSSPRGISLVGETLYIAAMHLF
 KKENGKIEKIDLSNSYEFINDIVNISGKTYLLAQNKEEELEVCELNKGDWTLKFKKPLKAYKFLKSVEEMA

t05A.aa BB009

CSSESIFSSQLGNLQKIKHEYNILGSSSPRGISLVGETLYIAAMHLFKKENGKIEKIDLSNSYEFINDIVNISGKTY
 LLAQNKEELEVCELNKGDWTLKFKKPLKAYKFLKSVEEMA

f06A.nt BB014

TAAGGAGCATATATGAGGATTGGTGGCGTTGTATAATAGCATTGGCTTATTGGTTGTATTGCGCTGATA
 ATCAGGAACAAGCTGTCAAACTTTTTGAGAATTGGAAAGTAGTGATATGGTTCCGATGAGATTGTTACTGA
 AGGCATATTCTAGTTAAATTATGCGTCTGAACATCGTTATTGGTGAGATAAAAAGACTTTAATTAGT
 TTTAAAGATCCTAATTATCNGNTGAGTACNCCCAGTGAGTGACTATAATGAGGAGTATTAAATTCTTTC
 TAGATTAGGGTCTGAGCAATCTAAAGACCTGATTAAGTTGTTATTATGGTAAAATGAGCAGAACATAATAA
 ATTATGCGTATAGTCGTTGGCTGTATTGAGTATAGAGGAGTTATATTCTCTAGATATTAGTATTCTGGCAG
 GGGAGCCATGAGTATAATCGTAATATGCCTAGACCCACTGCTTATGAACAATATTAAAAGTGAAGAGGTATGATT
 ATAATAGCCAGTTCTATTACCTACATAA

t06A.nt BB014

TGTTATTCGCCTGATAATCAGGAACAAGCTGTTCAAACCTTTTTGAGAATTGGAAAGTAGTGATATGGTTCCG
 ATGAGATTGTTACTGAAGGCATATTCTAGTTAAATTATGCGTCTGAACATCGTTATTGGTGAGATAAAA
 AAAGACTTTAATTAGTTAAAGATCCTAATTATCNGNTGAGTACNCCCAGTGAGTGACTATAATGAGGAGTAT
 TTTAATAAAATTCTTAGATTAGGGTCTGAGCAATCTAAAGACCTGATTAAGTTGTTATTATGGTAAAATG
 AGCAGAACATAATAAAATTATGCGTATAGTCGTTGGCTGTATTGAGTATAGAGGAGTTATATTCTCTAGATAT
 TAAGTATTCTGGCGAGGGGAGCCATGAGTATAATCGTAATATGCCTAGACCCACTGCTTATGAACAATATTAAA
 GTGAAGAGGTATGATTATAAT

f06A.aa BB014

GAYMRILVGCVIIALALLGCYLPDNQEQAQVQTFFENSESSDMGSDEIVTEGIFSSLKLYASEHRLLVEIKTLISL
 KDPNYXXVXPVSDYNEEYFNKFFLDLGSEQSKDLIKLFIMVKNEQNNNKFMRIVRWLYSCIEELYSLDIKYSGEG
 SHEYNRNMMPRTAYEQYLKVKRYDYNSPVSLPT

t06A.aa BB014

CYLPDNQEQAQVQTFFENSESSDMGSDEIVTEGIFSSLKLYASEHRLLVEIKTLISLKDPMYXXVXPVSDYNEEY
 FNKFFLDLGSEQSKDLIKLFIMVKNEQNNNKFMRIVRWLYSCIEELYSLDIKYSGEGSHEYNRNMMPRTAYEQYLK
 VKRYDYN

f07A.nt BB023

TAAAGTATTATTTTTTATTATCCACTGTTCTTGTCAAGAGACTGATGGATTAGCAGAGGCTTCTAAAA
 GGGCAGAGCCTGGAGAATTAGTTAGATTGCGAGCTTGTCAAGAGATCCAAGTCAACTAGACTGATCTTAC

TABLE 1. Nucleotide and Amino Acid Sequences

AAATTATGTTGATTATGTATATTGGCGCTTCTGGTATTGTTAAGCCGGAAGATATGGTTGAGATCTGGGATA
 AATAATTGGAGCGTTTACTTACTCCTTCTGCAAGGGTGCAGGCTTACGTTAAAATTCAAGTTGTTGCGCCCGCTG
 TTGTTAAGAGTGAGTCAAAAGGTACGCAGGTGATACTATTTAGGGTAAGAGTTGTTCCAAGCTATTCTCA
 ATCATCTGCTATGATTATGCCACCATTAAATTCCCTTTATTCAGGGAAAGTGGCAATCAATTAGGCAAA
 GGTCTTATTGATAACATTAAACCATGAAAGAAATTAAAGGTATCTGTTATAGTTAGGTATGAGATAGATCTG
 AGGTTTATTGAGATATGAATGNCATGGAATATGCTTNNCTATGGTACTTTAAAGTTAAAGGGTGGCTGA
 TTAAATTGTCAAATCCTAACTATATTCTAATATCATCCAGAATTAAAGACGATGTTCAAATTATCCT
 CTTGCTTCAAGTAAAATGAGATTAAAGGCTTTAGAGTTCAAAGTCACACAGTTCAAAGAGCAAATTCTCATCT
 TTATGTTAAAGATTAAAGAGTTCTTATGATAAGTTGAGTGTTCATAGATTCTGATATTGACAGTGAGTCTGT
 ATTAAAGTTATGAGACTAGCGGAAC TGAAATTAAAGGCAACAGNAACNTTAAAGNGTTTA
 AAGCTTAGAGAAAAAATTCTATGCCTGAAGGCTTTCCAAAACTTGAGAAAAGATTGAGAGTGAAAAACCTG
 AAGAATCATCTCCGAAAAATTAG

t07A.nt BB023

GAGGGTTCTAAAAGGGCAGAGCCTGGAGAATTAGTTAGATTTCGCCAGCTTGCAAGAGATCCAAGTTCAACTA
 GACTTGATCTTACAAATTATGTTGATTATGTATATTGGCGCTTCTGGTATTGTTAAGCCGGAAGATATGGTGT
 AGATCTTGGATAAAATAATTGGAGCGTTTACTTACTCCTTCTGCAAGGTGCAGGCTTACGTTAAAATTCAAGTT
 GTGCGCCCGCTGTTAAGAGTGAGTCAAAAGGTACGCAGGTGATACTATTTAGGGTAAGAGTTGTTTC
 CAAGCTATTCTCAATCATCTGCTATGATTATGCCACCATTTAAATTCCCTTTTATTCAAGGGAAAGTGGCAATCA
 ATTGTTAGGCAAAGGTCTTATTGATAACATTAAACCATGAAAGAAATTAAAGGTATCTGTTATAGTTAGGGTAT
 GAGATAGATCTGAGGTTTATTGAGATATGAATGNCATGGAATATGCTTNNCTATGGTACTTTAAAGTTA
 AAGGGTGGGCTGATTAAATTGGTCAAATTCTAACTATATTCTAATATATCATCCAGAATTAAAGACGATGT
 TCCAAATTATCCTCTTGCTTCAAGTAAAATGAGATTAAAGGCTTTAGAGTTCAAAGTCACACAGTTCAAAGAG
 CAAAATTCTATCTTTATGTTAAAGATTAAAGAGTTCTTATGATAAGTTGAGTGTTCATAGATTCTGATATTG
 ACAGTGAGTCTGTATTAAAGTTATGAGACTAGCGGAAC TGAAATTAAAGGCAACAGNAACNTT
 TAAAAGNGTTTAAAGCTTAGAGAAAAAATTCTATGCCTGAAGGCTTTCCAAAACTTGAGAAAAGATTGAG
 AGTAAAAACCTGAAGAATCATCTCCGAAAAAT

f07A.aa BB023

SILFFLLSTVLFAQETDGLAEGSKRAEPGELVLDFAELARDPSSTRLDLTNYVDYVYSGASGIVKPEDMVVDLGIN
 NWSVLLTPSARLQAYVKNSVVAAPAVVKSESKRYAGDTILGVRVLFPSYSQSSAMIMPPFKIPFYSGESGNQFLGKG
 LIDNIKTMKEIKVSVYSLGYEIDLEVLFEDMNXMNEYAXSMGTLFKGWADLIWSNPNYIPNISSRIIKDDVPNYPL
 ASSKMRFKAFRVSKSKEQNFIFYVKDLRVLYDKLVSIDSIDSEVFVYETSGTESLRKLKAHXTFKXVLK
 LREKISMPEGSFQNFVKEKPEESSPKN

t07A.aa BB023

EGSKRAEPGELVLDFAELARDPSSTRLDLTNYVDYVYSGASGIVKPEDMVVDLGINNWSVLLTPSARLQAYVKNSV
 VAPAVVKSESKRYAGDTILGVRVLFPSYSQSSAMIMPPFKIPFYSGESGNQFLGKLIDNIKTMKEIKVSVYSLGY
 EIDLEVLFEDMNXMNEYAXSMGTLFKGWADLIWSNPNYIPNISSRIIKDDVPNYPLASSKMRFKAFRVSKSKE
 QNFIFYVKDLRVLYDKLVSIDSIDSEVFVYETSGTESLRKLKAHXTFKXVLKREKISMPEGSFQNFVKEKIE
 SEKPEESSPKN

f08A.nt BB024

TGAATATTAATAATAAAAAAGGAGTAACAATGAAAATCATCAACATATTATTTGTTATTTTACTAATGCTAA
 ACGGCTGTAATTCTAATGATAATGACACTTAAAAACAAATGCCAACAAACAAAAAGACGGGAAAGCGTGATTT
 AACCCAAAAAGAAACAACACAAGAAAAACCAAATCTAAAGAAGAACTACTAGAGAAAAGCTATCTGACGATCAA
 AAAACACATCTTGACTGGTTAAAACCCGCTTAACTGGTGCTGGAGAATTGACAAATTCTAGAAAATGATGATG
 ATAAAATAAAATCAGCACTTGATCATATAAAACTCAACTTGATAGTTGATATGGTATCAAGCAGAACACAAAA
 AACCACTTCAAAACTGTGGTACAGAATTCTTAAAATGGTATAGATAATTGCAACTGGAGCGGTTAGT
 AACTGCAATAATGGTGGCTAA

t08A.nt BB024

TABLE 1. Nucleotide and Amino Acid Sequences

TGTAATTCTAATGATAATGACACTTAAAAAACAAATGCCAACAAACAAAAGACGGGGAAACCGTGATTAAACCC
 AAAAGAAACAAACACAAGAAAAACCAAAATCTAAAGAAGAACTACTTAGAGAAAAGCTATCTGACGATCAAAAAC
 ACATCTTGACTGGTTAAACCCGCTTAACTGGTGCTGGAGAATTGACAATTCTTAGAAAATGATGATGATAAA
 ATAAAATCAGCACTTGATCATATAAAACTCAACTTGATAGTTGTAATGGTATCAAGCAGAACACAAAAACCA
 CTTCAAAATGTGGTTACAGAATTCTTAAAATGGTATAGATAATTGCAACTGGAGCGGTTAGTAACTG
 CAATAATGGTGGC

f08A.aa BB024

ILIIKKGVTMKIINILFCLFLMLNGCNSNDNTLKNNQQTKRRGKRDLTQKETTQEKPKSKEELLREKLSDDQK
 THLDWLKPALTGAGEFDKFLENDDKIKSALDHKTQLDSCNGDQAEQQKTFKTVVTEFFKNGDIDNFATGAVSN
 CNNGG

t08A.aa BB024

CNSNDNTLKNNQQTKRRGKRDLS1TQKETTQEKPKSKEELLREKLSDDQKTHLDWLKPALTGAGEFDKFLENDD
 DKIKSALDHKTQLDSCNGDQAEQQKTFKTVVTEFFKNGDIDNFATGAVSNCNNGG

f09A.nt BB025

TGAATATTAAATAAAAAAGGAATAATAATGAAAATTATCACATATTATTTGTTTATTTTACTAATGCTAA
 ACGGCTGTAATTCTAATGATACTAATAATGCCAACAAAAGTAGACAAAACGTGATTTAACCCAAAAGAAC
 AACACAAGAAAAACCTAAATCTAAAGAAGAACTTCTTAGAGAAAAGCTAAATGATAATCAAAAACACACCTTGAC
 TGGTAAAAGAAGCTCTGGCAATGATGGAGAATTAAATAAAATTAGGATATGATGAAAGCAAAATAATCTG
 CACTGATCATATAAAGAGTGAACTTGACAGTTGACTGGAGATAAGGTTGAAAATAACCTCAAGCAGGT
 CGTCAGGAGGCCCTAAAGGGGGCATAGACGGCTTGAAAATACTGCAAGTAGTACGTGCAAAATTCAAA

t09A.nt BB025

TGTAATTCTAATGATACTAATAATGCCAACAAAAGTAGACAAAACGTGATTTAACCCAAAAGAAC
 AAGAAAAACCTAAATCTAAAGAAGAACTTCTTAGAGAAAAGCTAAATGATAATCAAAAACACACCTTGACTGGTT
 AAAAGAAGCTCTGGCAATGATGGAGAATTAAATAAAATTAGGATATGATGAAAGCAAAATAAAATCTGCACTT
 GATCATATAAAGAGTGAACTTGACAGTTGACTGGAGATAAGGTTGAAAATAACCTCAAGCAGGTGTT
 AGGAGGCCCTAAAGGGGGCATAGACGGCTTGAAAATACTGCAAGTAGTACGTGCAAAATTCA

f09A.aa BB025

ILIIKKGIIMKIINILFCLFLMLNGCNSNDTNNSQTKSRSRQKRDLTQKEATQEKPKSKEELLREKLNDNQKTHLDW
 LKEALGNDGEFNKFLGYDESKIKSALDHKSELDSCGDKVENKNTFKQVVQEALKGGIDGFENTASSTCKNS

t09A.aa BB025

CNSNDTNNSQTKSRSRQKRDLTQKEA51TQEKPKSKEELLREKLNDNQKTHLDWLKEALGNDGEFNKFLGYDESKIKS
 ALDHKSELDSCGDKVENKNTFKQVVQEALKGGIDGFENTASSTCKNS

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

Query	GenSeq Access No.	GenSeq Gene Description	BLAST Score	BLAST P-Value
f01A.aa	gi 2690256	(AE000790) antigen, P35, putative [Borrelia burgdorferi]	1523	5.90E-206
f02A.aa	gi 2690286	(AE000790) B. burgdorferi predicted coding region BBA69 [Borrelia	1320	2.10E-174
f02A.aa	gi 2690285	(AE000790) B. burgdorferi predicted coding region BBA68 [Borrelia	278	7.50E-71
f02A.aa	gi 2690105	(AE000789) B. burgdorferi predicted coding region BBI38 [Borrelia	151	8.40E-54
f02A.aa	gi 2690092	(AE000789) antigen, P35, putative [Borrelia burgdorferi]	151	2.70E-48
f02A.aa	gi 2690183	(AE000787) antigen, P35, putative [Borrelia burgdorferi]	155	4.20E-22
f02A.aa	gi 2690106	(AE000789) B. burgdorferi predicted coding region BBI39 [Borrelia	154	1.30E-21
f03A.aa	gi 2688051	(AE001127) antigen, S2, putative [Borrelia burgdorferi]	1223	7.60E-164
f03A.aa	gi 1063419	S2 gene product [Borrelia burgdorferi]	116	3.00E-22
f03A.aa	gi 2690227	(AE000790) antigen, S2 [Borrelia burgdorferi] >pir ID70207 D70207	116	9.70E-22
f03A.aa	gi 2690128	(AE000788) protein p23 [Borrelia burgdorferi] >pir C70257 C70257	110	5.70E-19
f03A.aa	gi 2689956	(AE000785) protein p23 [Borrelia burgdorferi] >pir D70225 D70225	104	7.90E-15
f04A.aa	gi 2690078	(AE000784) B. burgdorferi predicted coding region BBH18 [Borrelia	1873	5.60E-250
f04A.aa	gi 2690192	(AE000787) B. burgdorferi predicted coding region BBJ13 [Borrelia	167	1.40E-15
f05A.aa	gi 2687919	(AE001117) B. burgdorferi predicted coding region BB0028 [Borrelia	696	4.20E-92
f06A.aa	gi 2690129	(AE000788) outer membrane protein [Borrelia burgdorferi]	884	4.80E-124
f06A.aa	gi 2690089	(AE000789) conserved hypothetical protein [Borrelia burgdorferi]	731	2.20E-118
f06A.aa	gi 520783	unknown [Borrelia burgdorferi] >gi 551742 unknown [Borrelia	337	4.30E-58
f07A.aa	gi 2688608	(AE001168) flagellar filament outer layer protein (flaA) [Borrelia	1668	2.50E-224
f07A.aa	gi 1575447	FlaA protein [Borrelia burgdorferi] >gi 1019754 orf [Borrelia	1645	3.60E-221
f07A.aa	gi 152896	flagellar filament surface antigen [Spirochaeta aurantia]	144	1.70E-38
f07A.aa	gi 15059	endoflagellar sheath protein [Treponema pallidum]	139	3.80E-28
f07A.aa	gi 433524	flagellin FlaA1 [Serpulina hyodysenteriae] >gi 904393 endoflagellar	119	3.00E-26
f07A.aa	pir A32814	flagellar filament surface antigen - Spirochaeta aurantia A32814	116	9.40E-11
f08A.aa	gi 1209837	lipoprotein [Borrelia burgdorferi]	508	2.10E-78
f08A.aa	gi 2121280	(AF000270) lipoprotein [Borrelia burgdorferi] >gi 3095109	547	4.00E-70
f08A.aa	gi 1209873	lipoprotein [Borrelia burgdorferi]	303	3.70E-51

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f08A.aa	gil1209843	lipoprotein [Borrelia burgdorferi]	395	2.20E-49
f08A.aa	gil1209849	lipoprotein [Borrelia burgdorferi]	219	2.60E-27
f08A.aa	gil3095105	(AF046998) 2.9-8 lipoprotein [Borrelia burgdorferi]	234	4.30E-27
f08A.aa	gil1209831	lipoprotein [Borrelia burgdorferi]	209	1.10E-22
f08A.aa	gil3095107	(AF046999) 2.9-9 lipoprotein [Borrelia burgdorferi]	200	1.80E-22
f08A.aa	gil1209857	lipoprotein [Borrelia burgdorferi]	200	2.50E-21
f08A.aa	gnlIPDle26	surface-exposed lipoprotein [Borrelia afzelii] 8244	142	1.80E-11
f09A.aa	gil1209843	lipoprotein [Borrelia burgdorferi]	453	8.60E-67
f09A.aa	gil2121280	(AF000270) lipoprotein [Borrelia burgdorferi] >gil3095109	379	1.00E-56
f09A.aa	gil1209873	lipoprotein [Borrelia burgdorferi]	282	1.10E-45
f09A.aa	gil1209837	lipoprotein [Borrelia burgdorferi]	357	7.10E-44
f09A.aa	gil1209849	lipoprotein [Borrelia burgdorferi]	143	1.60E-13
f09A.aa	gnlIPDle26	surface-exposed lipoprotein [Borrelia afzelii] 8244	111	3.60E-13
f09A.aa	gil3095105	(AF046998) 2.9-8 lipoprotein [Borrelia burgdorferi]	142	5.40E-13
f101.aa	gil2688708	(AE001176) conserved hypothetical protein [Borrelia burgdorferi]	1099	4.50E-152
f105.aa	gil2688693	(AE001175) B. burgdorferi predicted coding region BB0758 [Borrelia	1276	2.20E-177
f11-12.aa	gil2690139	(AE000788) B. burgdorferi predicted coding region BBK01 [Borrelia	1473	4.70E-193
f11-12.aa	gil2690030	(AE000786) B. burgdorferi predicted coding region BBG01 [Borrelia	1066	1.40E-138
f11-12.aa	gil2690074	(AE000784) B. burgdorferi predicted coding region BBH37 [Borrelia	173	6.20E-93
f11-12.aa	gil2690188	(AE000787) B. burgdorferi predicted coding region BBJ08 [Borrelia	192	2.70E-75
f11-4.aa	gil2690150	(AE000788) B. burgdorferi predicted coding region BBK12 [Borrelia	1144	2.70E-147
f11-4.aa	gil2690145	(AE000788) B. burgdorferi predicted coding region BBK07 [Borrelia	852	5.70E-127
f11-4.aa	gil2690095	(AE000789) B. burgdorferi predicted coding region BB110 [Borrelia	153	1.30E-34
f11-4.aa	gil2690197	(AE000787) B. burgdorferi predicted coding region BBJ31 [Borrelia	115	1.40E-12
f11-4.aa	gil2690219	(AE000787) B. burgdorferi predicted coding region BBJ45 [Borrelia	115	1.40E-12
f112-1.aa	gil2690054	(AE000784) B. burgdorferi predicted coding region BBH06 [Borrelia	573	7.00E-75
f12.aa	gil2688785	(AE001182) B. burgdorferi predicted coding region BB0838 [Borrelia	6008	0
f129.aa	gil2688685	(AE001174) B. burgdorferi predicted coding region BB0739 [Borrelia	987	6.20E-133
f14-8.aa	gil2689955	(AE000785) antigen, P35, putative [Borrelia burgdorferi]	385	2.70E-75

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f14-8.aa	gi 2690120	(AE000789) <i>B. burgdorferi</i> predicted coding region BB134 [Borrelia	330	2.60E-66
f14-8.aa	gi 2690052	(AE000784) antigen, P35, putative [Borrelia burgdorferi]	287	4.00E-64
f14-8.aa	gi 2690100	(AE000789) <i>B. burgdorferi</i> predicted coding region BB116 [Borrelia	172	1.10E-38
f14-8.aa	gi 2690115	(AE000789) <i>B. burgdorferi</i> predicted coding region BB128 [Borrelia	173	1.70E-28
f14-8.aa	gi 2690116	(AE000789) <i>B. burgdorferi</i> predicted coding region BB129 [Borrelia	163	8.20E-24
f14-8.aa	gi 2690207	(AE000787) <i>B. burgdorferi</i> predicted coding region BB102 [Borrelia	220	1.90E-23
f14-8.aa	gi 2690099	(AE000789) <i>B. burgdorferi</i> predicted coding region BB115 [Borrelia	140	3.60E-12
f14-8.aa	gi 2690125	(AE000788) antigen, P35, putative [Borrelia burgdorferi]	111	1.00E-11
f142.aa	gi 2688655	(AE001172) glutamate transporter (gltP) [Borrelia burgdorferi]	2233	7.19999999999982e-311
f142.aa	gn IPIDe23	hypothetical protein [Bacillus subtilis] >gn IPIDe1182902	727	2.60E-156
f142.aa	gn IPIDd10	Proton/sodium-glutamate symport protein (Glutamate-aspartate	762	6.60E-146
f142.aa	gi 1574711	proton glutamate symport protein (gltP) [Haemophilus influenzae]	903	2.10E-131
f142.aa	gi 2983758	(AE000735) proton/sodium-glutamate symport protein [Aequifex	111	8.40E-36
f142.aa	gi 143000	proton glutamate symport protein [Bacillus stearothermophilus]	125	1.20E-30
f142.aa	gi 143002	proton glutamate symport protein [Bacillus caldotenax]	125	1.90E-28
f142.aa	gn IPIDe11	proton/sodium-glutamate symport protein [Bacillus subtilis]	122	2.20E-25
f142.aa	gn IPIDd10	glutamate transporter [Caenorhabditis elegans]	121	1.80E-22
	22697			
f142.aa	gi 1255318	coded for by <i>C. elegans</i> cDNA cm08h9; coded for by <i>C. elegans</i> cDNA	121	2.10E-22
f142.aa	gi 2388712	(AF017105) amino acid transporter [Chlamydia psittaci]	135	3.60E-22
f142.aa	gi 2655021	(AF018259) glutamate transporter 5A [Ambystoma tigrinum]	125	7.70E-22
f142.aa	gn IPIDe14	gltT-R gene product [Clostridium perfringens]	199	4.60E-21
f142.aa	gi 396412	gltP [Escherichia coli] >gi 147160 proton-glutamate [Escherichia	109	7.90E-21
f147.aa	gi 2688656	(AE001172) NADH oxidase, water-forming (nox) [Borrelia burgdorferi]	2245	7.20E-303
f147.aa	gi 642030	NADH oxidase [Serpulina hydysenteriae]	318	9.20E-105
f147.aa	gi 2650234	(AE001077) NADH oxidase (noxA-2) [Archaeoglobus fulgidus]	303	2.90E-93

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f147.aa	gi 2792490	(AF041467) coenzyme A disulfide reductase [Staphylococcus aureus]		194	2.60E-90
f147.aa	gi 2650383	(AE001088) NADH oxidase (noxA-1) [Archaeoglobus fulgidus]		286	3.30E-88
f147.aa	gn IPID1d10	H2O-forming NADH Oxidase [Streptococcus mutans] 09320		369	4.30E-85
f147.aa	gi 49023	NADH peroxidase [Enterococcus faecalis] >pir S18332 S18332 NADH		638	3.20E-83
f147.aa	gi 1591361	NADH oxidase (nox) [Methanococcus jannaschii] >pir A6438 I A6438I		535	4.80E-83
f147.aa	gi 2622461	(AE000898) NADH oxidase [Methanobacterium thermoautotrophicum]		303	8.40E-72
f147.aa	gi 47045	NADH oxidase [Enterococcus faecalis] >pir S26965 S26965 NADH oxidase		547	8.80E-71
f147.aa	gi 2650233	(AE001077) NADH oxidase (noxA-3) [Archaeoglobus fulgidus]		312	2.00E-63
f147.aa	gi 1674132	(AE000044) Mycoplasma pneumoniae, NADH oxidase; similar to		175	7.00E-61
f147.aa	gi 1045969	NADH oxidase [Mycoplasma genitalium] >pir D64230 D64230 NADH		164	4.10E-51
f147.aa	gi 2648692	(AE000975) NADH oxidase (noxA-5) [Archaeoglobus fulgidus]		143	2.00E-40
f147.aa	gi 2983379	(AE000709) NADH oxidase [Aquifex aeolicus]		162	5.50E-30
f150.aa	gi 2688659	(AE001172) conserved hypothetical protein [Borelia burgdorferi]		1319	2.70E-179
f150.aa	gi 2983887	(AE000743) hypothetical protein [Aquifex aeolicus]		238	1.40E-25
f150.aa	gi 2581796	(AF001974) putative TrkA [Thermoanaerobacter ethanolicus]		175	5.80E-23
f150.aa	gi 1377829	unknown [Bacillus subtilis] >gn IPID1 007628 orf4 [Bacillus subtilis]		212	1.50E-21
f150.aa	gn IPID1e11	similar to hypothetical proteins [Bacillus subtilis] 85982		181	6.00E-17
f150.aa	gn IPID1d10	hypothetical protein [Synechocystis sp.] >pir S75999 S75999 11497		128	3.70E-11
f152.aa	gi 2688660	(AE001172) K+ transport protein (mp) [Borelia burgdorferi]		2200	2.40000000 00 213e- 3 3
f152.aa	gi 2983882	(AE000743) K+ transport protein homolog [Aquifex aeolicus]		239	3.60E-106
f152.aa	gn IPID1e11	similar to Na+-transporting ATP synthase [Bacillus subtilis] 84940		158	6.60E-64
f152.aa	gn IPID1e11	similar to Na+-transporting ATP synthase [Bacillus subtilis] 85983		131	3.40E-62
f152.aa	gn IPID1d10	Na+-ATPase subunit J [Synechocystis sp.] >pir S75455 S75455 18749		141	1.70E-55

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f152.aa	gnl P D d10 Na+ -ATPase subunit J [Enterococcus hirae] 04799		209	4.00E-45
f152.aa	gi 2581795 [AF001974] putative TrkG [Thermoanaerobacter ethanolicus]		149	2.20E-29
f152.aa	gi 1674061 [AE000036] Mycoplasma pneumoniae, Na(+) translocating ATPase		104	4.00E-28
f152.aa	gi 1046024 Na+ -ATPase subunit J [Mycoplasma genitalium] >pir F64235 F64235 Na+		114	2.80E-27
f152.aa	gi 567062 HKT1 [Triticum aestivum] >pir S47582 S47582 high-affinity potassium		137	2.00E-17
f154.aa	gi 2688664 [AE001172] B. burgdorferi predicted coding region BB0722 [Borrelia	2456	0	
f157.aa	gi 2688641 [AE001171] rod shape-determining protein (mrkB-2) [Borrelia	2300	0	
f157.aa	gi 143657 endospore forming protein [Bacillus subtilis]		224	2.60E-61
f157.aa	gi 580938 internal open reading frame (AA 1-290) [Bacillus subtilis]		224	2.60E-61
f157.aa	gi 2982781 [AE000670] rod shape determining protein RodA [Aquifex aeolicus]	333	5.40E-61	
f157.aa	gi 580937 spoVE gene product (AA 1-366) [Bacillus subtilis] >gnl P D e1185111	224	7.70E-59	
f157.aa	gi 147695 rod-shape-determining protein [Escherichia coli] >gi 1778551	340	6.10E-58	
f157.aa	gnl P D e32 sfr [Streptomyces coelicolor] 8589		362	6.40E-58
f157.aa	gi 1572976 rod shape-determining protein (mrkB) [Haemophilus influenzae]	307	4.00E-56	
f157.aa	gnl P D e11 similar to cell-division protein [Bacillus subtilis] 85075	203	2.60E-45	
f157.aa	gi 1469784 putative cell division protein ftsW [Enterococcus hirae]	231	6.90E-45	
f157.aa	gi 1016213 strong sequence similarity to FtsW, RodA, and SpoV-E [Cyanophora	206	3.00E-41	
f157.aa	gnl P D d10 rod-shape-determining protein [Synechocystis sp.] 19002		184	1.60E-38
f157.aa	gi 146039 cell division protein [Escherichia coli] >gi 40857 FtsW protein		104	8.30E-35
f157.aa	gi 1574692 cell division protein (ftsW) [Haemophilus influenzae]		114	3.30E-33
f157.aa	gi 1165286 FtsW [Borrelia burgdorferi] >gi 2688164 (AE001137) cell division		170	6.20E-32
f17-6.aa	gi 2690100 [AE000789] B. burgdorferi predicted coding region BB116 [Borrelia	1250	1.70E-164	
f17-6.aa	gi 2690120 [AE000789] B. burgdorferi predicted coding region BB134 [Borrelia	142	3.40E-59	
f17-6.aa	gi 2690115 [AE000789] B. burgdorferi predicted coding region BB128 [Borrelia	447	6.70E-56	
f17-6.aa	gi 2690052 [AE000784] antigen, P35, putative [Borrelia burgdorferi]		182	1.10E-34
f17-6.aa	gi 2689955 [AE000785] antigen, P35, putative [Borrelia burgdorferi]		196	6.60E-34

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f17-6.aa	gi 2690114	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI27 [Borrelia	176	1.00E-16
f17-6.aa	gn IPD 010 12343	gene required for phosphorylation of oligosaccharides/ has	178	2.80E-15
f17-6.aa	gi 2690207	(AE000787) <i>B. burgdorferi</i> predicted coding region BBJ02 [Borrelia	114	3.50E-13
f17-6.aa	gn IPD 32 9895	(AJ000496) cyclic nucleotide-gated channel beta subunit	152	1.10E-11
f170.aa	gi 2688652	(AE001171) <i>B. burgdorferi</i> predicted coding region BB0708 [Borrelia	524	2.60E-73
f186.aa	gi 2688622	(AE001169) <i>B. burgdorferi</i> predicted coding region BB0689 [Borrelia	792	1.80E-105
f186.aa	gi 2688622	(AE001169) <i>B. burgdorferi</i> predicted coding region BB0689 [Borrelia	792	1.80E-105
f19-2.aa	gi 2690120	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI34 [Borrelia	1341	2.70E-177
f19-2.aa	gi 2689955	(AE000785) antigen, P35, putative [Borrelia burgdorferi]	347	7.00E-53
f19-2.aa	gi 2690052	(AE000784) antigen, P35, putative [Borrelia burgdorferi]	254	7.70E-53
f19-2.aa	gi 2690100	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI16 [Borrelia	142	6.60E-50
f19-2.aa	gi 2690115	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI28 [Borrelia	144	7.60E-34
f19-2.aa	gi 2690116	(AE000789) <i>B. burgdorferi</i> predicted coding region BB129 [Borrelia	183	2.20E-21
f19-2.aa	gi 2690207	(AE000787) <i>B. burgdorferi</i> predicted coding region BBJ02 [Borrelia	171	2.00E-16
f19-2.aa	gi 2690099	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI15 [Borrelia	166	1.20E-15
f19-2.aa	gi 2690125	(AE000788) antigen, P35, putative [Borrelia burgdorferi]	122	5.70E-14
f19-4.aa	gi 2690116	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI29 [Borrelia	1129	1.30E-150
f19-4.aa	gi 2690099	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI15 [Borrelia	260	3.00E-30
f19-4.aa	gi 2689955	(AE000785) antigen, P35, putative [Borrelia burgdorferi]	180	1.80E-23
f19-4.aa	gi 2690120	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI34 [Borrelia	183	1.50E-21
f19-4.aa	gi 2690052	(AE000784) antigen, P35, putative [Borrelia burgdorferi]	192	1.20E-19
f19-4.aa	gi 2690207	(AE000787) <i>B. burgdorferi</i> predicted coding region BBJ02 [Borrelia	149	8.90E-14
f19-4.aa	gi 2690098	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI14 [Borrelia	138	8.00E-12
f19-6.aa	gi 2690115	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI28 [Borrelia	995	1.20E-131
f19-6.aa	gi 2690100	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI16 [Borrelia	447	3.00E-55
f19-6.aa	gi 2689955	(AE000785) antigen, P35, putative [Borrelia burgdorferi]	219	2.00E-36
f19-6.aa	gi 2690120	(AE000789) <i>B. burgdorferi</i> predicted coding region BBI34 [Borrelia	144	3.50E-34
f19-6.aa	gi 2690052	(AE000784) antigen, P35, putative [Borrelia burgdorferi]	130	6.30E-12
f196.aa	gi 2688620	(AE001169) methyl-accepting chemotaxis protein (mcp-5) [Borrelia	3093	0

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f196_aa	gi 2688621	(AE001169) methyl-accepting chemotaxis protein (mcp-4) [Borrelia	615	1.90E-83
f196_aa	gi 496484	lipC gene product [Bacillus subtilis] >pir 40496 40496 methylation	180	6.90E-28
f196_aa	gnl PDI d10	methyl-accepting chemotaxis protein TlpC [Bacillus subtilis]	180	4.90E-27
f196_aa	gnl PDI d11	methyl-accepting chemotaxis protein [Bacillus subtilis]	162	5.10E-25
	73493			
f196_aa	gi 882594	ORF f506 [Escherichia coli] >gi 1789453 (AE000389) aerotaxis	204	1.70E-24
f196_aa	gi 148350	tas [Enterobacter aerogenes] >pir D32302 D32302 probable aspartate	179	1.80E-24
f196_aa	gi 1066850	putative [Rhodobacter capsulatus] >pir JC4735 JC4735	207	1.80E-24
f196_aa	gi 154381	chemoreceptor [Salmonella typhimurium] >pir A47178 A47178	230	2.00E-24
f196_aa	gi 459690	transmembrane receptor [Bacillus subtilis] >gnl PDI d1185997	212	1.40E-23
f196_aa	gi 805015	MCpA protein [Rhodobacter sphaeroides] >pir S70094 S54262	237	2.10E-23
f196_aa	gi 40424	mcpA gene product [Caulobacter crescentus] >pir S23064 S23064 mcpa	238	7.30E-23
f196_aa	gi 144913	sensory transducer protein [Clostridium thermocellum]	227	8.90E-23
f196_aa	gi 1061063	Trg sensory transducer protein [Escherichia coli]	211	2.40E-20
f196_aa	gnl PDI d10	Methyl-accepting chemotaxis protein III (MCP-III) (Ribose and	211	2.50E-20
f197_aa	gi 2688621	(AE001169) methyl-accepting chemotaxis protein (mcp-4) [Borrelia	3724	0
	gi 2688620	(AE001169) methyl-accepting chemotaxis protein (mcp-5) [Borrelia	615	8.40E-83
f197_aa	gi 1066850	putative [Rhodobacter capsulatus] >pir JC4735 JC4735	227	9.80E-27
f197_aa	gi 882594	ORF f506 [Escherichia coli] >gi 1789453 (AE000389) aerotaxis	217	1.00E-26
f197_aa	gi 154381	chemoreceptor [Salmonella typhimurium] >pir A47178 A47178	239	2.80E-25
f197_aa	gi 496484	lipC gene product [Bacillus subtilis] >pir 40496 40496 methylation	202	5.10E-25
f197_aa	gnl PDI d10	methyl-accepting chemotaxis protein TlpC [Bacillus subtilis]	202	5.10E-25
f197_aa	gi 2564665	(AF022807) putative methyl accepting chemotaxis protein [Rhizobium	212	7.20E-24
	gi 459691	transmembrane receptor [Bacillus subtilis] >gnl PDI d1185996	215	1.10E-23
f197_aa	gi 43218	serine chemoreceptor [Escherichia coli] >bbsl127562 serine	236	2.80E-23
f197_aa	gi 537197	CG Site No. 63; alternate gene name cheD [Escherichia coli]	236	2.90E-23
f197_aa	gi 448077	methyl-accepting chemotaxis protein I [Escherichia coli] >gi 2367378	236	2.90E-23
f197_aa	gnl PDI d10	transducer [Pseudomonas aeruginosa]	178	4.20E-23

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f197_aa	gj148349	tsr [Enterobacter aerogenes] >pir[C32302 C32302] serine transducer	234	5.50E-23
f197_aa	gi 2626835	chemotactic transducer [Pseudomonas aeruginosa]	177	5.70E-23
f200_aa	gi 2688600	(AE001168) ribose/galactose ABC transporter, permease protein	1887	5.10E-266
f200_aa	gn IPD 31	unknown [Bacillus subtilis] >gn IPD 31 184234 similar to 1453	283	1.50E-63
f200_aa	gi 2649711	(AE001042) ribose ABC transporter, permease protein (rbsC-1)	202	1.10E-47
f200_aa	gi 2130609	(AF000308) putative polytopic protein [Mycoplasma fermentans]	119	2.10E-27
f200_aa	gn IPD 31	unknown [Bacillus subtilis] >gn IPD 31 184235 similar to 1493	112	1.10E-18
f200_aa	gi 950073	membrane forming protein [Mycoplasma capricolum] >pir S77790 S77790	161	5.60E-16
f200_aa	gi 2688599	(AE001168) ribose/galactose ABC transporter, permease protein	108	2.00E-14
f208_aa	gi 2688610	(AE001168) B. burgdorferi predicted coding region BB0674 [Borrelia	1726	6.70E-244
f21-4_aa	gi 197833	Bbk2.11 [Borrelia burgdorferi] >pir S70531 S70531 bbk2.11 protein	474	3.00E-70
f21-4_aa	gi 2627267	ErpL [Borrelia burgdorferi]	477	6.30E-69
f21-4_aa	gi 1707281	putative outer membrane protein [Borrelia burgdorferi]	503	6.60E-66
f21-4_aa	gi 896042	CspF [Borrelia burgdorferi] >pir S70532 S70532 outer surface protein	503	6.60E-66
f21-4_aa	gi 1707287	putative outer membrane protein [Borrelia burgdorferi]	489	3.00E-60
f21-4_aa	gi 1707290	putative outer surface protein [Borrelia burgdorferi]	342	3.20E-49
f21-4_aa	gi 163633	ErpK [Borrelia burgdorferi]	268	1.70E-48
f21-4_aa	gi 466482	outer surface protein F [Borrelia burgdorferi] >pir I40287 I40287	321	3.80E-38
f21-4_aa	gi 896038	Bbk2.10 precursor [Borrelia burgdorferi] >pir S70534 S70534 bbk2.10	121	3.90E-34
f21-4_aa	gi 896040	Bbk2.10 precursor [Borrelia burgdorferi] >pir S70533 S70533 bbk2.10	118	2.30E-33
f21-4_aa	gi 051120	outer surface protein G [Borrelia burgdorferi] >gi 1373118 ErpG	107	3.30E-33
f21-4_aa	gi 2444428	(AF020657) ErpX protein [Borrelia burgdorferi]	118	6.00E-14
f210_aa	gi 2688603	(AE001168) conserved hypothetical protein [Borrelia burgdorferi]	867	2.60E-116
f210_aa	gi 2688604	(AE001168) chemotaxis response regulator (cheY-3) [Borrelia	733	1.40E-97
f210_aa	gi 408274	CheY [Borrelia burgdorferi]	720	9.00E-96
f210_aa	gi 1765976	chemotaxis protein CheY [Treponema pallidum]	405	6.60E-52
f210_aa	gi 142682	chemotactic response protein [Bacillus subtilis] >gn IPD 31 185224	184	8.00E-30
f210_aa	gi 940149	CheY [Thermotoga maritima]	171	1.50E-27

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f210.aa	gil2649557	(AE001031) chemotaxis response regulator (cheY) [Archaeoglobus	168	1.50E-26
f210.aa	gil620085	cheY gene product [Listeria monocytogenes]	183	3.00E-26
f210.aa	gnlIPDle24	YneI [Bacillus subtilis] >gil870926 response regulator	166	4.00E-24
f210.aa	9646			
f210.aa	gil149620	ORF2 [Leptospira borgpetersenii] >sp P24086 YLB3_LEPIN	121	4.70E-22
		HYPOTHETICAL		
f210.aa	gil1408275	orfX; putative OrfX protein [Borrelia burgdorferi]	208	9.20E-22
f210.aa	gil1994802	cheY gene product [Halobacterium salinarium] >pir S58645 S58645_CheY	139	8.90E-18
f210.aa	gil143598	spoOF [Bacillus subtilis] >gil143601 SpoOF protein [Bacillus	113	4.70E-11
f216.aa	gil2688586	(AE001167) conserved hypothetical protein [Borrelia burgdorferi]	804	1.20E-109
f216.aa	gil1575446	orfA [Borrelia burgdorferi]	472	1.10E-91
f219.aa	gil2688594	(AE001167) B. burgdorferi predicted coding region BB0664 [Borrelia	1122	3.10E-148
f22.aa	gil2688779	(AE001181) B. burgdorferi predicted coding region BB0832 [Borrelia	1400	4.90E-188
f22.aa	gil2688779	(AE001181) B. burgdorferi predicted coding region BB0832 [Borrelia	1400	4.90E-188
f221.aa	gil2688596	(AE001167) B. burgdorferi predicted coding region BB0662 [Borrelia	692	2.60E-93
f229.aa	gil2688591	(AE001167) oxygen-independent coproporphyrinogen III oxidase,	863	7.80E-120
f24-1.aa	gil2039285	putative vls recombination cassette Vls6 [Borrelia burgdorferi]	924	1.80E-114
f24-1.aa	gil2039284	putative vls recombination cassette Vls5 [Borrelia burgdorferi]	867	6.30E-107
f24-1.aa	gil2039287	putative vls recombination cassette Vls8 [Borrelia burgdorferi]	824	1.50E-104
f24-1.aa	gil2039289	putative vls recombination cassette Vls10 [Borrelia burgdorferi]	829	7.50E-102
f24-1.aa	gil2039320	vmp-like sequence protein VlsE [Borrelia burgdorferi]	644	1.10E-98
f24-1.aa	gil2039288	putative vls recombination cassette Vls9 [Borrelia burgdorferi]	783	8.20E-96
f24-1.aa	gil2039330	vmp-like sequence protein VlsE [Borrelia burgdorferi]	742	6.30E-95
f24-1.aa	gil2039336	vmp-like sequence protein VlsE [Borrelia burgdorferi]	509	1.50E-92
f24-1.aa	gil2039286	putative vls recombination cassette Vls7 [Borrelia burgdorferi]	754	6.60E-92
f24-1.aa	gil2039324	vmp-like sequence protein VlsE [Borrelia burgdorferi]	488	8.10E-86
f24-1.aa	gil2039316	vmp-like sequence protein VlsE [Borrelia burgdorferi]	531	1.70E-85
f24-1.aa	gil2039312	vmp-like sequence protein VlsE [Borrelia burgdorferi]	531	1.20E-83
f24-1.aa	gil2039326	vmp-like sequence protein VlsE [Borrelia burgdorferi]	476	2.00E-82
f24-1.aa	gil2039332	vmp-like sequence protein VlsE [Borrelia burgdorferi]	474	5.10E-82
f24-1.aa	gil2039328	vmp-like sequence protein VlsE [Borrelia burgdorferi]	420	3.50E-59

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f253_aa	gi 2688567 [AE001165] Na+/H+ antiporter (nhaC-1) [Borrelia burgdorferi]	2247	0
f253_aa	gi 2688566 [AE001165] Na+/H+ antiporter (nhaC-2) [Borrelia burgdorferi]	609	6.40E-155
f253_aa	gi 2209268 Na+/H+ antiporter [Bacillus firmus] >pir A41594 A41594	158	9.40E-15
f253_aa	gi 1574661 Na+/H+ antiporter (nhaC) [Haemophilus influenzae]	143	4.20E-14
f253_aa	gnl P D e11 similar to Na+/H+ antiporter [Bacillus subtilis] 85625	137	1.20E-11
f253_aa	gnl P D e32 hypothetical protein [Bacillus subtilis] >gnl P D e1182969 4972	133	2.00E-11
f265_aa	gi 2688555 [AE001164] conserved hypothetical protein [Borrelia burgdorferi]	1196	9.90E-161
f269_aa	gi 2688560 [AE001164] B. burgdorferi predicted coding region BB0624 [Borrelia	1654	5.50E-226
f28_2_aa	gi 2690174 [AE0007788] B. burgdorferi predicted coding region BBK47 [Borrelia	1683	2.80E-222
f28_2_aa	gi 2690161 [AE0007788] B. burgdorferi predicted coding region BBK49 [Borrelia	1068	2.20E-163
f28_3_aa	gi 2690138 [AE0007788] immunogenic protein P37, putative [Borrelia burgdorferi]	281	6.00E-48
f28_3_aa	gi 2690127 [AE0007788] immunogenic protein P37 [Borrelia burgdorferi]	209	3.20E-28
f28_3_aa	gi 2459605 immunogenic protein P37 [Borrelia burgdorferi]	208	4.50E-28
f28_3_aa	gi 2690137 [AE0007788] immunogenic protein P37, putative [Borrelia burgdorferi]	172	5.50E-17
f29_aa	gi 2688764 [AE001180] B. burgdorferi predicted coding region BB0826 [Borrelia	869	8.20E-116
f290_aa	gi 2688537 [AE001162] serine-type D-Ala-D-Ala carboxypeptidase (dacA)	2046	1.50E-281
f290_aa	gi 143439 DD-carboxypeptidase [Bacillus subtilis] >pir B42708 B42708	161	6.60E-36
f290_aa	gnl P D e11 D-alanyl-D-alanine carboxypeptidase (penicillin binding 85617	161	6.60E-36
f290_aa	gnl P D d10 Probable penicillin-binding protein. [Escherichia coli] 16562	131	3.30E-28
f290_aa	sp P37604 DACD_SA LTY PENICILLIN-BINDING PROTEIN 6B PRECURSOR	135	9.10E-28
f290_aa	gi 1572974 penicillin-binding protein 5 (dacA) [Haemophilus influenzae]	145	3.00E-27
f290_aa	gi 580849 D-alanine carboxypeptidase [Bacillus stearothermophilus]	170	4.10E-27
f290_aa	gi 1778549 penicillin-binding protein 5 [Escherichia coli] >gi 41212 precursor	152	3.20E-26
f290_aa	gi 142820 penicillin-binding protein 5 [Bacillus subtilis]	137	4.60E-26
f290_aa	gi 410134 penicillin-binding protein [Bacillus subtilis] >gnl P D e1185588	137	4.60E-26
f290_aa	gi 41218 precursor [Escherichia coli]	136	1.30E-25

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f290.aa	gnlPDIid10	Penicillin-binding protein 6 precursor (D-alanyl-D-alanine)		136	1.30E-25
f290.aa	gil1864022	penicillin binding protein 4 [Staphylococcus aureus]		155	5.10E-22
f290.aa	gnlPDIid15	penicillin binding protein 4 [Staphylococcus aureus]		155	5.10E-22
f290.aa	4145				
f290.aa	gnlPDIid26	penicillin-binding protein 4 [Staphylococcus aureus]		155	5.10E-22
f290.aa	46682				
f291.aa	gil2688538	(AE001162) L-lactate permease (lctP) [Borrelia burgdorferi]	2473	0	
f291.aa	gnlPDIid27	lactate permease [Streptococcus iniae]	586	1.20E-132	
f291.aa	4704				
f291.aa	gil882504	ORF f560 [Escherichia coli] >gil1789347 (AE000380) f560; This 560 aa	345	3.60E-95	
f291.aa	gil2313225	(AE000535) L-lactate permease (lctP) [Helicobacter pylori]	359	1.10E-94	
f291.aa	gil2313224	(AE000535) L-lactate permease (lctP) [Helicobacter pylori]	348	2.90E-93	
f291.aa	gil404693	L-lactate permease [Escherichia coli] >gil466741 aug is 3rd start	331	7.20E-82	
f291.aa	gnlPDIid31	hypothetical protein [Bacillus subtilis] >gnlPDIid1186107	330	9.00E-80	
f291.aa	3006				
f291.aa	gnlPDIid10	lactate permease [Bacillus subtilis]	300	1.70E-61	
f291.aa	22632				
f291.aa	gnlPDIid11	L-lactate permease [Bacillus subtilis] >pirF69649F69649	300	1.10E-60	
f291.aa	82258				
f291.aa	gnlPDIid10	homologue of L-lactate permease of E. coli [Bacillus	265	6.40E-56	
f291.aa	09575				
f291.aa	gil2649804	(AE001049) L-lactate permease (lctP) [Archaeoglobus fulgidus]	170	1.50E-47	
f291.aa	gnlPDIid28	L-lactate permease [Sulfolobus solfataricus]	163	2.60E-44	
f291.aa	3914				
f291.aa	gil1574148	L-lactate permease (lctP) [Haemophilus influenzae]	173	6.00E-35	
f296.aa	gil2688517	(AE001161) chaperonin, putative [Borrelia burgdorferi]	1276	4.40E-177	
f296.aa	gil840643	mucZ gene product [Coxiella burnetii] >pirI40852I40852 mucZ	101	7.90E-12	
f3.aa	gil26888797	(AE001183) B. burgdorferi predicted coding region BB0844 [Borrelia	1604	1.40E-211	
f30.aa	gil26888765	(AE001180) B. burgdorferi predicted coding region BB0825 [Borrelia	1343	2.00E-181	
f301.aa	gil2688521	(AE001161) methyl-accepting chemotaxis protein (mcp-3) [Borrelia	2756	0	
f301.aa	gil1805311	methyl-accepting chemotaxis protein B [Treponema denticola]	211	7.00E-20	

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f301.aa	gi 2688522	[AE001161] methyl-accepting chemotaxis protein (mcp-2) [Borrelia	189	2.80E-18
f301.aa	gi 2367665	[AF016689] Mcp-2 [Treponema pallidum]	189	3.50E-17
f301.aa	gi 2352917	[AF012922] methyl-accepting chemotaxis protein [Treponema	187	5.70E-17
f301.aa	gi 1354776	[MCP-1 [Treponema pallidum]	189	5.90E-17
f301.aa	gi 2619023	[AF027868] YoaH [Bacillus subtilis] >gnlP Dle1185333 similar to	184	2.80E-16
f301.aa	gi 1654421	transducer HtrB protein [Halobacterium salinarum]	177	2.20E-15
f301.aa	gi 415694	chemoreceptor [Desulfovibrio vulgaris] >pir G36943 G36943	163	3.50E-15
f301.aa	gi 459691	transmembrane receptor [Bacillus subtilis] >gnlP Dle1185996	163	4.90E-15
f301.aa	gi 2104730	ORF2 [Desulfurococcus sp. SY]	173	5.80E-15
f301.aa	gi 2914132	methyl accepting chemotaxis homolog [Treponema denticola]	170	1.10E-14
f301.aa	gi 459689	transmembrane receptor [Bacillus subtilis] >gnlP Dle1185998	164	1.30E-14
f301.aa	gi 496484	lipC gene product [Bacillus subtilis] >pir I40496 I40496 methylation	170	3.80E-14
f301.aa	gi 2313163	(AE000530) methyl-accepting chemotaxis transducer (lpC)	170	6.30E-14
f308.aa	gi 2688527	(AE001161) B. burgdorferi predicted coding region BB0592 [Borrelia	1227	1.70E-176
f31-2.aa	gi 2690202	(AE000787) B. burgdorferi predicted coding region BBJ36 [Borrelia	1771	7.20E-235
f31-2.aa	gi 2690200	(AE000787) B. burgdorferi predicted coding region BBJ34 [Borrelia	423	4.60E-88
f31.aa	gi 2688766	(AE001180) B. burgdorferi predicted coding region BB0824 [Borrelia	957	7.80E-133
f314.aa	gi 2688509	(AE001160) pfs protein (pfs-2) [Borrelia burgdorferi]	1329	7.40E-180
f314.aa	gi 2690087	(AE000789) pfs protein (pfs) [Borrelia burgdorferi]	335	1.50E-77
f314.aa	gi 2688288	(AE001143) pfs protein (pfs-1) [Borrelia burgdorferi]	266	1.00E-65
f314.aa	gi 2738591	(AF012886) Pfs [Buchnera aphidicola]	115	1.70E-52
f314.aa	gi 1552737	similar to purine nucleoside phosphorylase (deoD) [Escherichia	133	6.90E-52
f314.aa	gnlP Dle11	similar to purine nucleoside phosphorylase [Bacillus	157	1.20E-49
	83957			
f314.aa	gi 147158	pfs [Escherichia coli] >gi 457107 ORF [Escherichia coli] [SUB 9-219]	133	2.50E-42
f314.aa	gi 1574146	pfs protein (pfs) [Haemophilus influenzae] >pir C64169 C64169 pfs	110	2.70E-37
f314.aa	gi 2267164	(AF009177) pfs protein homolog [Helicobacter pylori]	118	3.30E-23
f314.aa	gi 2313168	(AE000530) pfs protein (pfs) [Helicobacter pylori]	115	1.00E-22
f314.aa	gi 777939	Pfs [Treponema pallidum]	102	1.90E-20
f314.aa	gi 2689970	(AE000785) B. burgdorferi predicted coding region BBE07 [Borrelia	191	1.50E-19
f314.aa	gnlP Dle24	unknown [Mycobacterium tuberculosis] >sp Q10889 Y05A_MYCTU	105	7.60E-16

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9405				
f32-4_aa	gil2690221 (AE0007787) B. burgdorferi predicted coding region BBY47 [Borrelia	1192	4.00E-163	
f32-4_aa	gil2689979 (AE0007785) B. burgdorferi predicted coding region BBE16 [Borrelia	103	4.10E-11	
f32_aa	gil2688767 (AE0011180) B. burgdorferi predicted coding region BB0823 [Borrelia	623	1.80E-81	
f32_aa	gil2688767 (AE0011180) B. burgdorferi predicted coding region BB0823 [Borrelia	623	1.80E-81	
f320_aa	gil2688497 (AE0011159) carboxypeptidase, putative [Borrelia burgdorferi]	1373	6.40E-186	
f320_aa	gil2529473 (AF0066665) YokZ [Bacillus subtilis]	136	9.80E-28	
f320_aa	gil2415396 (AF015775) carboxypeptidase [Bacillus subtilis] >gnl PfD 1185433	136	1.90E-27	
f320_aa	gil1209528 D,D-carboxypeptidase [Enterococcus faecalis]	148	3.30E-16	
	>spl Q47746 VANY ENTFA			
f320_aa	gil155044 vanY [Transposon Tn1546] >gil149126 D,D-carboxypeptidase [Plasmid	142	1.60E-13	
f328_aa	gil2688502 (AE0011159) CTP synthase (pyrG) [Borrelia burgdorferi]	869	6.10E-119	
f328_aa	gil1591801 CTP synthase (pyrG) [Methanococcus annaschii] >pir E64446 E64446	325	6.20E-59	
f328_aa	gil2650385 (AE001088) CTP synthase (pyrG) [Archaeoglobus fulgidus]	304	4.20E-54	
f328_aa	gil1399854 CTP synthetase [Synechococcus PCC7942] >sp Q54775 PYRG_SYN P7	313	3.30E-52	
f328_aa	gnl PfD 10 CTP synthetase [Synechocystis sp.] >pir S75840 S75840 CTP	295	1.80E-50	
	19032			
f328_aa	gil143597 CTP synthetase [Bacillus subtilis] >gil 853762 CTP synthase [Bacillus	274	1.60E-49	
f328_aa	gil2983754 (AE000735) CTP synthetase [Aequifex aeolicus]	271	1.50E-46	
f328_aa	gil1574630 CTP synthetase (pyrG) [Haemophilus influenzae] >pir F64181 F64181	234	1.90E-44	
f328_aa	gil413755 CTP synthetase [Spiroplasma citri] >sp P52200 PYRG_SPICI CTP	231	3.00E-44	
f328_aa	gil2621483 (AE000826) CTP synthase [Methanobacterium thermoautotrophicum]	257	2.80E-40	
f328_aa	gil950067 CTP synthase [Mycoplasma capricolum] >pir S77767 S77767 CTP	220	4.10E-39	
	synthase			
f328_aa	gil904007 cytidine triphosphate synthetase precursor [Giardia intestinalis]	219	2.00E-38	
f328_aa	gil147478 CTP synthetase (EC 6.3.4.2) [Escherichia coli]	217	2.90E-38	
f328_aa	gil882674 CTP synthetase [Escherichia coli] >gil 789142 (AE000361) CTP	214	7.70E-38	
f328_aa	gil38688 CTP synthase [Azospirillum brasilense] >pir I39496 S25101 CTP	132	3.20E-37	
f342_aa	gil2688495 (AE0011158) B. burgdorferi predicted coding region BB0563 [Borrelia	944	5.30E-130	
f346_aa	gil1272356 phosphotransferase enzyme II [Borrelia burgdorferi] >gil2688474	828	1.10E-108	

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f346.aa	gil145603	PTS enzyme III glc [Escherichia coli] >gil145605 PTS enzyme III glc	385	8.80E-53
f346.aa	gil1314675	glucose-specific component IIIA of the PTS system [Escherichia coli]	385	9.30E-53
f346.aa	gil47658	III(Glc) (cr) (AA 1 - 169) [Salmonella typhimurium]	382	2.30E-52
f346.aa	gil1574566	glucose phosphotransferase enzyme III-glc (cr) [Haemophilus	397	8.70E-50
f346.aa	gil143819	nagE gene product [Klebsiella pneumoniae] >pir186071S18607	349	2.80E-41
f346.aa	gil146913	N-acetylglucosamine transport protein [Escherichia coli]	334	3.20E-39
f346.aa	gil1072418	glcA [Staphylococcus carnosus] >pir1S46952S46952	317	7.20E-37
f346.aa	gil1072419	glcB [Staphylococcus carnosus] >pir1S636061S46953	315	1.40E-36
f346.aa	gil1146177	phosphotransferase system glucose-specific enzyme II [Bacillus	295	7.30E-36
f346.aa	gil529001	PTS glucose-specific permease [Bacillus stearothermophilus]	294	8.80E-36
f346.aa	gnlIPIDle11	alternate gene name: yzfA; similar to phosphotransferase	293	1.40E-33
	82187			
f346.aa	gil580912	enzyme III-glucose [Bacillus subtilis]	257	1.20E-30
f346.aa	gil602681	phosphocarrier protein (enzyme IIa) [Mycoplasma capricolum]	243	1.00E-28
f346.aa	gil1432153	cellulose-specific PTS permease [Klebsiella oxytoca]	257	1.20E-28
f352.aa	gil2688482	(AE001157) B. burgdorferi predicted coding region BB0553 [Borrelia	2547	0
f352.aa	gil2688482	(AE001157) B. burgdorferi predicted coding region BB0553 [Borrelia	1005	1.30E-132
f363.aa	gil2688468	(AE001156) B. burgdorferi predicted coding region BB0543 [Borrelia	1109	5.40E-153
f368.aa	gil2688450	(AE001155) conserved hypothetical integral membrane protein	1133	4.10E-157
f368.aa	gil1787004	(AE000181) o234; This 234 aa ORF is 26 pct identical (15 gaps) to	417	1.40E-67
f368.aa	gil2314055	(AE000601) conserved hypothetical integral membrane protein	129	3.50E-16
f368.aa	gnlIPIDle12	S1R [Cowpox virus]	135	1.80E-14
	89272			
f368.aa	gnlIPIDd10	24K membrane protein [Pseudomonas aeruginosa]	108	9.00E-13
	03176			
f368.aa	gil41284	put. 23.5-kd protein [Escherichia coli] >gil1787205 (AE000199)	101	1.00E-11
f371.aa	gil2688452	(AE001155) conserved hypothetical protein [Borrelia burgdorferi]	1066	3.60E-143
f371.aa	gil2196997	Orf256 [Treponema pallidum]	154	1.10E-15
f373.aa	gil2688453	(AE001155) zinc protease, putative [Borrelia burgdorferi]	3663	0
f373.aa	gil1574200	hypothetical [Haemophilus influenzae] >pir1E64171IE64171	295	2.70E-67
f373.aa	gil1787770	(AE000246) f931; residues 5-650 are 99 pct identical to YDDC_ECOLI	289	1.10E-57

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f373.aa	gi 535004	ccs106 gene product [Escherichia coli]		289	3.20E-57
f373.aa	gi 799369	metalloendopeptidase [Pisum sativum]		148	7.10E-28
f373.aa	gi 2827039	(AF008444) chloroplast processing enzyme [Arabidopsis thaliana]		150	1.70E-26
f373.aa	gi 2983709	(AE000732) processing protease [Aquifex aeolicus]		136	4.30E-24
f373.aa	gi 2314155	(AE000609) protease (pqqE) [Helicobacter pylori] >pir D64646 D64646		115	5.30E-23
f378.aa	gi 2688458	(AE001155) B. burgdorferi predicted coding region BB0531 [Borrelia		1030	1.30E-136
f384.aa	gi 2688435	(AE001154) inositol monophosphatase [Borrelia burgdorferi]		1470	3.80E-201
f4-15.aa	gi 2690238	(AE000790) surface lipoprotein P27 [Borrelia burgdorferi]		1400	1.50E-185
f4-15.aa	gi 144008	P27 [Borrelia burgdorferi] >pir S34995 S34995 surface lipoprotein		462	2.40E-96
f4-50.aa	gi 2690243	(AE000790) decorin binding protein B (dbpB) [Borrelia burgdorferi]		900	6.30E-117
f4-50.aa	gi 2062381	decorin binding protein B [Borrelia burgdorferi]		897	1.60E-116
f4-50.aa	gi 2809217	(AF042796) putative decorin-binding protein precursor [Borrelia		887	3.60E-115
f4-50.aa	gi 2809218	(AF042796) decorin-binding protein precursor [Borrelia burgdorferi]		172	2.00E-33
f4-50.aa	gi 2690249	(AE000790) decorin binding protein A (dbpA) [Borrelia burgdorferi]		176	9.50E-33
f4-50.aa	gi 2062379	decorin binding protein A [Borrelia burgdorferi]		177	6.10E-32
f4-66.aa	gi 2690229	(AE000790) chpA1 protein, putative [Borrelia burgdorferi]		807	1.60E-107
f4.aa	gi 2688787	(AE001183) conserved hypothetical integral membrane protein		2408	0
f4.aa	gi 2697115	(AF008219) unknown [Borrelia afzelii]		1138	1.90E-305
f4.aa	gi 1573583	H. influenzae predicted coding region H10594 [Haemophilus		337	2.10E-109
f4.aa	gi 1788636	(AE000319) 0513; This 513 aa ORF is 31 pct identical (30 gaps) to		327	9.10E-80
f4.aa	gnl P D d10	homologue of hypothetical protein H10594 of H. influenzae		357	5.40E-69
f42-1.aa	gi 2689993	(AE000794) conserved hypothetical protein [Borrelia burgdorferi]		495	2.70E-62
f42-1.aa	gi 2689934	(AE000793) conserved hypothetical protein [Borrelia burgdorferi]		312	1.00E-37
f43-3.aa	gi 1209843	lipoprotein [Borrelia burgdorferi]		546	1.50E-69
f43-3.aa	gi 2121280	(AF000270) lipoprotein [Borrelia burgdorferi] >gi 3095109		442	1.80E-55
f43-3.aa	gi 1209837	lipoprotein [Borrelia burgdorferi]		365	3.10E-55
f43-3.aa	gi 1209873	lipoprotein [Borrelia burgdorferi]		269	5.30E-32
f43-3.aa	gi 1209849	lipoprotein [Borrelia burgdorferi]		141	1.70E-13
f43-3.aa	gi 3095105	(AF046998) 2.9-8 lipoprotein [Borrelia burgdorferi]		140	9.60E-13
f43-3.aa	gi 3095107	(AF046999) 2.9-9 lipoprotein [Borrelia burgdorferi]		132	1.40E-11

TABLE 2. Closest matching sequences between the poly peptides of the present invention and sequences in GenBank and Derwent databases.

f43.aa	gi 2688752	(AE001179) B. burgdorferi predicted coding region BB0811 [Borrelia	2337	6.60000000
			084856e-	
			315	
f446.aa	gi 2688383	(AE001151) B. burgdorferi predicted coding region BB0464 [Borrelia	920	7.20E-124
f45-2.aa	gi 1699017	ErpB2 [Borrelia burgdorferi] >gi 1373133 ErpB [Borrelia	364	7.50E-78
f45-2.aa	gi 2627270	ErpJ [Borrelia burgdorferi]	364	2.50E-77
f45-2.aa	gi 2627268	ErpM [Borrelia burgdorferi]	452	9.70E-60
f45-2.aa	gi 1373144	ErpD [Borrelia burgdorferi]	316	1.60E-58
f45-2.aa	gi 2444428	(AF020657) ErpX protein [Borrelia burgdorferi]	380	2.80E-55
f45-2.aa	gi 1051120	outer surface protein G [Borrelia burgdorferi] >gi 1373118 ErpG	213	7.10E-35
f45-2.aa	gi 1663633	ErpK [Borrelia burgdorferi]	152	1.60E-21
f45-2.aa	gn P D e32	(AJ000496) cyclic nucleotide-gated channel beta subunit	198	2.80E-16
	9895			
f45-2.aa	gi 466482	outer surface protein F [Borrelia burgdorferi] >pir 40287 40287	111	5.70E-14
f45-2.aa	gi 2246532	ORF 73, contains large complex repeat CR 73 [Kaposi's	174	5.90E-14
f45-2.aa	gi 160299	glutamic acid-rich protein [Plasmodium falciparum]	169	1.00E-13
f45-2.aa	gi 1707287	putative outer membrane protein [Borrelia burgdorferi]	101	2.20E-13
f45-2.aa	gi 1633572	Herpesvirus saimiri ORF73 homolog [Kaposi's sarcoma-associated	175	4.10E-13
f45-2.aa	gn P D d10	gene required for phosphorylation of oligosaccharides/ has	166	5.60E-13
	12343			
f45-2.aa	gi 2690100	(AE000789) B. burgdorferi predicted coding region BB116 [Borrelia	161	2.70E-12
f457 aa	gi 2688369	(AE001150) B. burgdorferi predicted coding region BB0456 [Borrelia	1021	6.20E-139
f469.aa	gi 2688368	(AE001150) Na+/H+ antiporter (napA) [Borrelia burgdorferi]	1544	1.10E-211
f47-2.aa	gi 1209849	lipoprotein [Borrelia burgdorferi]	742	2.30E-97
f47-2.aa	gi 1209857	lipoprotein [Borrelia burgdorferi]	407	7.80E-86
f47-2.aa	gi 1209831	lipoprotein [Borrelia burgdorferi]	393	5.00E-82
f47-2.aa	gn P D e26	surface-exposed lipoprotein [Borrelia burgdorferi]	321	2.60E-73
	8245			
f47-2.aa	gi 1209874	lipoprotein [Borrelia burgdorferi]	348	1.10E-64
f47-2.aa	gn P D e26	surface-exposed lipoprotein [Borrelia garinii]	333	1.40E-57
	8239			
f47-2.aa	gn P D e26	surface-exposed lipoprotein [Borrelia afzelii]	292	9.60E-44

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f47-2.aa	8244	gi 3095107 (AF046999) 2.9-9 lipoprotein [Borrelia burgdorferi]	328	3.80E-40
f47-2.aa	gn IPD e26	surface-exposed lipoprotein [Borrelia garinii]	320	1.70E-39
f47-2.aa	8242			
f47-2.aa	gi 1209837	lipoprotein [Borrelia burgdorferi]	210	4.80E-29
f47-2.aa	gi 2121280	(AF000270) lipoprotein [Borrelia burgdorferi] >gi 3095109	205	1.10E-27
f47-2.aa	gi 3095105	(AF046998) 2.9-8 lipoprotein [Borrelia burgdorferi]	217	6.30E-25
f47-2.aa	gi 1209873	lipoprotein [Borrelia burgdorferi]	113	2.40E-11
f477.aa	gi 2688350	(AE001149) fructose-bisphosphate aldolase (fba) [Borrelia	1506	3.60E-202
f477.aa	gi 82454	fructose 1,6-bisphosphate aldolase [Escherichia coli] >gi 41423	651	1.10E-131
f477.aa	gi 2708661	(AF037440) fructose 1,6-bisphosphate aldolase [Edwardsiella	593	1.40E-124
f477.aa	gi 1573507	fructose-bisphosphate aldolase (fba) [Haemophilus influenzae]	560	8.50E-120
f477.aa	gi 671841	fructose 1,6-bisphosphate aldolase [Campylobacter jejuni]	856	3.80E-113
f477.aa	gn IPD d10	fructose 1,6-bisphosphate aldolase [Schizosaccharomyces	749	1.70E-98
f477.aa	04756			
f477.aa	gi 433637	yeast fructose-bisphate-aldolase [Saccharomyces cerevisiae] >gi 3696	459	1.20E-92
f477.aa	gn IPD e19	fructose-1,6-bisphosphate aldolase [Euglena gracilis]	701	6.30E-92
f477.aa	0134			
f477.aa	gi 1334980	fructose 1,6 bisphosphate-aldolase [Neurospora crassa]	647	1.50E-84
f477.aa	gi 40495	fructose-bisphosphate aldolase [Corynebacterium glutamicum]	204	6.80E-37
f477.aa	gn IPD e31	Fba [Mycobacterium tuberculosis]	207	1.50E-35
f477.aa	5480			
f477.aa	gi 1045692	fructose-bisphosphate aldolase [Mycoplasma genitalium]	108	2.10E-23
f477.aa	gn IPD d10	hypothetical protein [Bacillus subtilis] >gn IPD e1184692	102	2.70E-15
f477.aa	03809			
f488.aa	gi 2688338	(AE001148) DNA gyrase, subunit A (gyrA) [Borrelia burgdorferi]	3222	0
f488.aa	gi 1790876	DNA gyrase subunit A [Clostridium acetobutylicum]	822	1.80E-171
f488.aa	gi 2650163	(AE001072) DNA gyrase, subunit A (gyrA) [Archaeoglobus fulgidus]	483	1.10E-162
f488.aa	gi 40019	ORF 821 (aa 1-821) [Bacillus subtilis] >gn IPD d1005785 A subunit of	836	6.10E-159
f488.aa	gi 459929	gyrase A subunit [Pseudomonas aeruginosa] >sp P48372 GYRA_PSEAE	418	7.00E-155
f488.aa	DNA			
f488.aa	gi 144206	DNA gyrase A [Campylobacter jejuni] >pir A48902 A48902 DNA gyrase	508	7.50E-154

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f488.aa	gi 466275	gyrase A [Mycobacterium tuberculosis] >sp Q07702 GYRA_MYCTU	395	3.50E-152
	DNA			
f488.aa	gn IPDle26	GyrA [Mycobacterium tuberculosis]	395	2.00E-151
	6924			
f488.aa	gi 43485	DNA gyrase A subunit [Haloferax] >pir S30571 S30571 DNA topoisomerase	275	6.10E-151
		(AB010081) A subunit of DNA gyrase [Bacillus sp.]		
f488.aa	gn IPDld10		549	1.20E-150
	25098			
f488.aa	gi IPDle21	DNA gyrase subunit A [Mycobacterium smegmatis]	388	5.90E-150
	4031			
f488.aa	gi 2731385	DNA gyrase [Serratia marcescens]	378	6.00E-148
f488.aa	gn IPDle13	DNA topoisomerase (ATP-hydrolysing) [Mycobacterium smegmatis]	388	7.30E-147
	7038			
f488.aa	gi 41634	gyrA gene product (AA 1-875) [Escherichia coli] >gi 41636 DNA gyrase	383	2.40E-146
		DNA gyrase subunit A [Mycoplasma genitalium]		
f488.aa	gi 497648		514	5.20E-146
		putative vls recombination cassette Vls3 [Borrelia burgdorferi]		
f49-2.aa	gi 2039282		943	2.30E-120
f49-2.aa	gi 2547241	vmp-like sequence protein VlSE [Borrelia burgdorferi]	434	4.10E-106
f49-2.aa	gi 2039324	vmp-like sequence protein VlSE [Borrelia burgdorferi]	458	3.00E-104
f49-2.aa	gi 2039281	putative vls recombination cassette Vls2 [Borrelia burgdorferi]	793	1.80E-100
f49-2.aa	gi 2039283	putative vls recombination cassette Vls4 [Borrelia burgdorferi]	729	4.60E-92
f49-2.aa	gi 2039308	vmp-like sequence protein VlSE [Borrelia burgdorferi]	652	1.40E-88
f49-2.aa	gi 2039288	putative vls recombination cassette Vls9 [Borrelia burgdorferi]	352	1.80E-88
f49-2.aa	gi 2039332	vmp-like sequence protein VlSE [Borrelia burgdorferi]	550	4.40E-88
f49-2.aa	gi 2039328	vmp-like sequence protein VlSE [Borrelia burgdorferi]	629	1.50E-85
f49-2.aa	gi 2039336	vmp-like sequence protein VlSE [Borrelia burgdorferi]	460	1.40E-82
f49-2.aa	gi 2039318	vmp-like sequence protein VlSE [Borrelia burgdorferi]	367	6.20E-82
f49-2.aa	gi 2039320	vmp-like sequence protein VlSE [Borrelia burgdorferi]	449	1.80E-77
f49-2.aa	gi 2483796	VlsE1 [Borrelia burgdorferi]	497	8.20E-76
f49-2.aa	gi 2039326	vmp-like sequence protein VlSE [Borrelia burgdorferi]	427	2.50E-64
f49-2.aa	gi 2039291	putative vls recombination cassette Vls13 [Borrelia burgdorferi]	409	1.30E-47
f494.aa	gi 2688346	(AE001148) B. burgdorferi predicted coding region BB0428 [Borrelia	547	8.20E-74
f5-14.aa	gi 2627268	ErpM [Borrelia burgdorferi]	1836	2.60E-236

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f5-14_aa	gi 1373144	ErpD [Borrelia burgdorferi]		543	4.40E-87
f5-14_aa	gi 2627270	ErpJ [Borrelia burgdorferi]		503	4.30E-83
f5-14_aa	gi 1699017	ErpB2 [Borrelia burgdorferi] >gi 1373133 ErpB [Borrelia burgdorferi]		503	2.60E-82
f5-14_aa	gi 2444428	(AF020657) ErpX protein [Borrelia burgdorferi]		399	9.30E-57
f5-14_aa	gnl PDIe32 9895	(AJ000496) cyclic nucleotide-gated channel beta subunit		228	1.50E-20
f5-14_aa	gnl PDId10 12343	gene required for phosphorylation of oligosaccharides/has		203	8.70E-18
f5-14_aa	gi 2246532	ORF 73, contains large complex repeat CR 73 [Kaposi's		197	3.30E-17
f5-14_aa	gi 1633572	Herpesvirus saimiri ORF73 homolog [Kaposi's sarcoma-associated		192	1.20E-16
f5-14_aa	gi 3068583	(AF000580) Rep-like [Dictyostelium discoideum]		197	3.60E-16
f5-14_aa	gi 2690100	(AE000789) B. burgdorferi predicted coding region BBII16 [Borrelia		183	2.90E-15
f5-14_aa	gi 1825739	No definition line found [Caenorhabditis elegans]		168	1.60E-14
f5-14_aa	gi 3044185	(AF056936) mature parasite-infected erythrocyte surface antigen		166	2.00E-14
f5-14_aa	gnl PDIe34 9084	E02A10.2 [Caenorhabditis elegans]		176	2.30E-14
f5-14_aa	gi 1051120	outer surface protein G [Borrelia burgdorferi] >gi 1373118 ErpG		157	3.30E-12
f5-15_aa	gi 2627267	ErpL [Borrelia burgdorferi]		1152	4.40E-147
f5-15_aa	gi 1197833	Bpk2.11 [Borrelia burgdorferi] >pir S705311S70531 bbk2.11 protein		856	3.30E-108
f5-15_aa	gi 896042	OspF [Borrelia burgdorferi] >pir S705321S70532 outer surface protein		325	1.00E-72
f5-15_aa	gi 1707281	putative outer membrane protein [Borrelia burgdorferi]		323	1.80E-72
f5-15_aa	gi 1707287	putative outer membrane protein [Borrelia burgdorferi]		322	6.60E-70
f5-15_aa	gi 466482	outer surface protein F [Borrelia burgdorferi] >pir 40287 40287		448	6.80E-68
f5-15_aa	gi 1707290	putative outer surface protein [Borrelia burgdorferi]		290	1.90E-52
f5-15_aa	gi 1663633	ErpK [Borrelia burgdorferi]		172	8.70E-43
f5-15_aa	gi 896038	Bpk2.10 precursor [Borrelia burgdorferi] >pir S705341S70534 bbk2.10		153	1.10E-42
f5-15_aa	gi 896040	Bpk2.10 precursor [Borrelia burgdorferi] >pir S705331S70533 bbk2.10		124	4.30E-39
f5-15_aa	gi 1051120	outer surface protein G [Borrelia burgdorferi] >gi 1373118 ErpG		105	3.10E-23
f5-15_aa	gi 1373144	ErpD [Borrelia burgdorferi]		103	1.10E-14
f50_aa	gi 2688754	(AE001179) B. burgdorferi predicted coding region BB0806 [Borrelia		2651	0
f502_aa	gi 26888313	(AE001146) sensory transduction histidine kinase, putative		7570	0

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f502_aa	gnlIPIDid10 25877	(AB006363) homologue of histidine kinase [Candida albicans]		296	3.80E-58
f502_aa	gi 354473	Os-1p [Neurospora crassa]		275	3.30E-57
f502_aa	gi 1679757	two-component histidine kinase CHK-1 [Glomerella cingulata]		382	4.20E-57
f502_aa	gi 1262208	Nik-1 [Neurospora crassa] >gi 1262210 Nik-1 [Neurospora crassa]		273	6.30E-57
f502_aa	gi 2460283	(AF024654) hybrid histidine kinase DHKB [Dictyostelium discoideum]		273	3.90E-55
f502_aa	gnlIPIDid10 17789	sensory transduction histidine kinase [Synechocystis sp.]		288	8.50E-54
f502_aa	gi 2623815	(AF030352) two component sensor [Pseudomonas aeruginosa]		252	4.00E-52
f502_aa	gi 939724	putative sensor kinase, regulatory protein for production of		252	1.80E-50
f502_aa	gi 151329	regulatory protein [Pseudomonas syringae] >sp P48027 LEM1_PSESY		248	1.20E-49
f502_aa	gi B41863 B41863	two-component regulatory protein lemA - Pseudomonas syringae		248	1.30E-49
f502_aa	gnlIPIDid10 18725	sensory transduction histidine kinase [Synechocystis sp.]		252	2.10E-49
f502_aa	gnlIPIDid10 02185	sensor-regulator protein [Escherichia coli] >gi 1789149		262	6.20E-49
f502_aa	gi 463195	pectate lyase [Pseudomonas viridisflava]		247	7.50E-49
f502_aa	gnlIPIDid10 18731	sensory transduction histidine kinase [Synechocystis sp.]		244	1.00E-48
f51-2_aa	gi 2444428	(AF020657) ErpX protein [Borrelia burgdorferi]		1755	2.20E-227
f51-2_aa	gi 2627268	ErpM [Borrelia burgdorferi]		399	3.20E-57
f51-2_aa	gi 373144	ErpD [Borrelia burgdorferi]		282	2.20E-50
f51-2_aa	gi 2627270	ErpJ [Borrelia burgdorferi]		271	6.00E-54
f51-2_aa	gi 699017	ErpB2 [Borrelia burgdorferi] >gi 1373133 ErpB [Borrelia		271	2.50E-33
f51-2_aa	gi 1051120	outer surface protein G [Borrelia burgdorferi] >gi 1373118 ErpG		109	3.70E-22
f51-2_aa	gnlIPIDid10 12343	gene required for phosphorylation of oligosaccharides/ has		203	5.40E-18
f51-2_aa	gi 1707287	putative outer membrane protein [Borrelia burgdorferi]		111	7.50E-18
f51-2_aa	gi 896042	OspF [Borrelia burgdorferi] >sp S70532 S70532 outer surface protein		111	2.10E-17
f51-2_aa	gi 1707281	putative outer membrane protein [Borrelia burgdorferi]		111	7.50E-17
f51-2_aa	gnlIPDle32 (AJ000496)	cyclic nucleotide-gated channel beta subunit		198	1.60E-16

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

	9895					
f51-2.aa	gi 2246532	ORF 73, contains large complex repeat CR 73 [Kaposi's			176	2.30E-14
f51-2.aa	gnl PTD e34	E02A10.2 [Caenorhabditis elegans]			170	2.10E-13
f51-2.aa	gi 160299	glutamic acid-rich protein [Plasmodium falciparum]			157	7.30E-12
f516.aa	gi 2688326	(AE001146) B. burgdorferi predicted coding region BB0409 [Borrelia			1096	2.00E-150
f517.aa	gi 2688320	(AE001146) PTS system, fructose-specific IIABC component (fruA-1)			1637	2.30E-228
f517.aa	gnl PTD e11	similar to fructose phosphotransferase system enzyme II			256	4.00E-88
f517.aa	gi 396296	similar to phosphotransferase system enzyme II [Escherichia coli]			305	9.10E-86
f517.aa	gi 405893	fructose-specific IIBC component [Escherichia coli] >gi 450372			224	4.30E-84
f517.aa	gi 151932	fructose enzyme II [Rhodobacter capsulatus] >gi 46021 fructose			222	4.70E-79
f517.aa	gi 1573422	fructose-permease IIBC component (fruA) [Haemophilus influenzae]			225	6.90E-69
f517.aa	gi 2688554	(AE001164) PTS system, fructose-specific IIABC component (fruA-2)			236	8.20E-66
f517.aa	gnl PTD e11	phosphotransferase system (PTS) fructose-specific enzyme IIBC			195	2.80E-65
	85030					
f517.aa	gi 155369	PTS enzyme-II fructose [Xanthomonas campestris] >pir B40944 B40944			187	8.10E-62
f517.aa	gi 305003	similar to fructose-specific phosphotransferase enzyme II			145	1.90E-39
f517.aa	gnl PTD d10	HsA [Escherichia coli] >gi 1786951 (AE000176)			148	2.80E-39
	11544					
f517.aa	gi 1813488	phosphotransferase enzyme II [Bacillus firmus]			226	3.90E-39
f517.aa	gi 757734	fruA gene product [Bacillus amyloliquefaciens] >pir S59965 S59965			177	2.50E-36
f517.aa	gnl PTD d10	PTS SYSTEM, FRUCTOSE-SPECIFIC IIBC COMPONENT (EIIBC-FRU)			173	1.10E-34
f517.aa	gi 1673731	(AE000010) Mycoplasma pneumoniae, fructose-permease IIBC component			143	9.00E-33
f519.aa	gi 2688327	(AE001146) B. burgdorferi predicted coding region BB0406 [Borrelia			1060	5.70E-145
f519.aa	gi 2688328	(AE001146) B. burgdorferi predicted coding region BB0405 [Borrelia			261	1.20E-47
f520.aa	gi 2688328	(AE001146) B. burgdorferi predicted coding region BB0405 [Borrelia			1022	3.90E-138
f520.aa	gi 2688327	(AE001146) B. burgdorferi predicted coding region BB0406 [Borrelia			261	4.00E-47
f523.aa	gi 2688300	(AE001145) glutamate transporter, putative [Borrelia burgdorferi]			2007	9.90E-284
f526.aa	gi 2688309	(AE001145) B. burgdorferi predicted coding region BB0399 [Borrelia			1087	1.60E-145

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f527.aa	gi 2688310 (AE001145) B. burgdorferi predicted coding region	BB0398 [Borrelia	1814	7.60E-249
f541.aa	gi 508421 antigen P39 [Borrelia burgdorferi] >gi 2688281 (AE001143) basic		1706	5.40E-230
f541.aa	gi 1753225 BmpA protein [Borrelia burgdorferi]		1698	6.80E-229
f541.aa	gn IPDle11 bmpA(p39,ORF1) [Borrelia burgdorferi]		1695	1.70E-228
f541.aa	gn IPDle11 membrane protein A [Borrelia burgdorferi] >gi 516592 membrane		1642	3.40E-221
f541.aa	gn IPDle11 membrane protein A [Borrelia burgdorferi]		1638	1.20E-220
f541.aa	gn IPDle11 bmpA(p39,ORF1) [Borrelia burgdorferi]		1551	1.00E-208
f541.aa	gn IPDle11 membrane protein A [Borrelia afzelii]		1502	5.60E-202
f541.aa	gn IPDle11 membrane protein A [Borrelia afzelii]		1499	1.40E-201
f541.aa	gn IPDle11 membrane protein A [Borrelia garinii]		1496	3.70E-201
f541.aa	gn IPDle11 membrane protein A [Borrelia afzelii]		1493	9.60E-201
f541.aa	gn IPDle11 membrane protein A [Borrelia garinii]		1488	4.60E-200
f541.aa	gn IPDle23 membrane protein A [Borrelia garinii]		1216	1.20E-162
f541.aa	gn IPDle23 membrane protein A [Borrelia garinii]		1211	5.90E-162
f541.aa	gn IPDle23 membrane protein A [Borrelia garinii]		1098	2.00E-146
f541.aa	gi 2688282 (AE001143) basic membrane protein B (bmpB) [Borrelia burgdorferi]		518	1.20E-123
f542.aa	gi 508422 [Borrelia burgdorferi immunodominant antigen P39 gene, complete		711	1.70E-95
f542.aa	gi 2688282 (AE001143) basic membrane protein B (bmpB) [Borrelia burgdorferi]		711	1.70E-95
f542.aa	gi 551744 membrane lipoprotein [Borrelia burgdorferi]		708	8.60E-95
f542.aa	gn IPDle11 bmpB(p39,ORF2) [Borrelia burgdorferi]		699	8.20E-94

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f542_aa	72836	gnlIPIDle11 72832	bmpB(p39,ORF2) [Borrelia afzelii]		634	1.00E-84
f542_aa	72839	gnlIPIDle11 7209	bmpB(p39,ORF2) [Borrelia garinii]		613	9.20E-82
f542_aa	72828	gnlIPIDle23 7214	membrane protein A [Borrelia garinii]		153	1.70E-32
f542_aa	72828	gnlIPIDle11 7214	bmpA(p39,ORF1) [Borrelia burgdorferi]		144	3.80E-32
f542_aa	72825	gnlIPIDle11 72833	membrane protein A [Borrelia garinii]		153	2.00E-31
f542_aa	72833	gnlIPIDle11 72837	BmpA protein [Borrelia burgdorferi]		155	2.80E-31
f542_aa	72833	gnlIPIDle11 72837	bmpA(p39,ORF1) [Borrelia burgdorferi]		155	2.80E-31
f542_aa	728421	gnlIPIDle11 72837	antigen P39 [Borrelia burgdorferi] >gi 2688281 (AE001143) basic		155	2.80E-31
f542_aa	72829	gnlIPIDle11 72830	membrane protein A [Borrelia garinii]		156	1.00E-30
f542_aa	72829	gnlIPIDle11 72830	membrane protein A [Borrelia afzelii]		144	1.90E-30
f542_aa	72829	gnlIPIDle11 72830	membrane protein A [Borrelia afzelii]		144	2.70E-30
f544_aa	72688284	gi 2688284 (AE001143) Mg2+ transport protein (mgtE) [Borrelia burgdorferi]			860	4.20E-119
f544_aa	72532228	gi 7532228 MgtE [Borrelia burgdorferi]			855	2.20E-118
f544_aa	72619724	gi 619724 MgtE [Bacillus firmus] >gi 40201140201 mgtE protein - Bacillus			176	3.70E-37
f544_aa	72802882	gi 7802882 extended ORF of mgtE gene; transcription from this start point is			182	1.30E-34
f544_aa	5479	gnlIPIDle31 5479	unknown [Mycobacterium tuberculosis]		183	4.50E-31
f544_aa	18132	gnlIPIDle10 18132	Mg2+ transporter [Synechocystis sp.] >pir S77552 S77552 Mg2+		165	4.60E-31
f544_aa	81529	gnlIPIDle11 81529	(AJ002571) YkoK [Bacillus subtilis] >gnlIPIDle1183350 similar		142	2.30E-30
f544_aa	72621701	gi 2621701 (AE000843) Mg2+ transporter [Methanobacterium thermoautotrophicum]			142	3.20E-21

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f545_aa	gi 2688284 (AE001143) Mg2+ transport protein (mgtE) [Borrelia burgdorferi]	860	4.20E-119
f545_aa	gi 1753228 MgtE [Borrelia burgdorferi]	855	2.20E-118
f545_aa	gi 619724 MgtE [Bacillus firmus] >pir 4020 140201 mgtE protein - Bacillus	176	3.70E-37
f545_aa	gi 780282 extended ORF of mgtE gene; transcription from this start point is	182	1.30E-34
f545_aa	gnl PDIe31 unknown [Mycobacterium tuberculosis]	183	4.50E-31
f545_aa	gnl PDIe31 15479 Mg2+ transporter [Synechocystis sp.] >pir S77552 S77552 Mg2+	165	4.60E-31
f545_aa	gnl PDIe11 18132 (AJ002571) YkoK [Bacillus subtilis] >gnl PDIe1183350 similar	142	2.30E-30
f545_aa	gi 2621701 (AE000843) Mg2+ transporter [Methanobacterium thermoautotrophicum]	142	3.20E-21
f561_aa	gi 49245 lipoprotein [Borrelia burgdorferi] >gi 2688271 (AE001142) lipoprotein	1000	1.30E-132
f561_aa	gi 495738 P22 [Borrelia burgdorferi]	982	3.70E-130
f577_aa	gi 2688261 (AE001141) B. burgdorferi predicted coding region BB0352 [Borrelia	1930	4.00E-264
f584_aa	gi 2688246 (AE001140) B. burgdorferi predicted coding region BB0346 [Borrelia	1094	4.10E-147
f596_aa	gi 2688241 (AE001140) P26 [Borrelia burgdorferi] >pir G7014 1G70141 P26	1322	1.20E-180
f596_aa	gi 2281465 (AF000366) P26 [Borrelia burgdorferi] >gi 2281465 (AF000366) P26	1010	5.90E-137
f598_aa	gi 2281462 (AF000366) oligopeptide permease homolog D [Borrelia burgdorferi]	652	1.20E-85
f598_aa	gi 143607 sporulation protein [Bacillus subtilis]	372	1.20E-45
f598_aa	gnl PDIe11 83166 oligopeptide ABC transporter (ATP-binding protein) [Bacillus	372	1.20E-45
f598_aa	gi 1574676 oligopeptide transport ATP-binding protein (oppD) [Haemophilus	344	6.70E-42
f598_aa	gi 677943 AppD [Bacillus subtilis] >gnl PDIe1183156 oligopeptide ABC	344	8.00E-42
f598_aa	gi 1787051 (AE000185) o612; 48 pct identical (33 gaps) to 525 residues from	346	2.50E-41
f598_aa	gi 47346 AmiE protein [Streptococcus pneumoniae] >pir S11152 S11152 amiE	338	1.10E-40
f598_aa	gi 47805 Opp D (AA1-335) [Salmonella typhimurium] >sp P04285 OPPD_SALTY	332	5.70E-40
f598_aa	pir A03413 oligopeptide transport protein oppD - Salmonella typhimurium	332	5.70E-40
f598_aa	gi 1787499 (AE000223) oligopeptide transport ATP-binding protein OppD	332	5.90E-40
f598_aa	gnl PDIe10 15494 Oligopeptide transport ATP-binding protein OppD. [Escherichia	332	5.90E-40
f598_aa	gi 495177 ATP binding protein [Lactococcus lactis] >sp P50980 OPPD_LACLC	331	8.40E-40

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f598_aa	gnlPDIe18	oligopeptide permease [Streptococcus pyogenes]	331	1.10E-39
f598_aa	7587	ATP binding protein [Lactococcus lactis] >pirlA53290lA53290	329	1.60E-39
f598_aa	gi 308850	ATP binding protein [Lactococcus lactis] >pirlA53290lA53290	322	2.30E-39
f598_aa	gi 2313399	(AE000548) dipeptide ABC transporter, ATP-binding protein (dppD)	565	4.30E-73
f6-21_aa	gi 2281468	(AF000948) OppAIV [Borrelia burgdorferi] >gi 2689891 (AE000792)	315	1.20E-37
f6-21_aa	gi 2253286	(AF005657) plasminogen binding protein [Borrelia burgdorferi]	314	1.60E-37
f6-21_aa	gi 2688228	(AE001139) oligopeptide ABC transporter, periplasmic	314	1.60E-37
f6-21_aa	gi 2809544	(AF043071) oligopeptide permease periplasmic binding protein	314	1.60E-37
f6-21_aa	gi 2281457	(AF000366) oligopeptide permease homolog AII [Borrelia burgdorferi]	314	1.60E-37
f6-21_aa	gi 2688227	(AE001139) oligopeptide ABC transporter, periplasmic	290	3.90E-34
f6-21_aa	gi 2281458	(AF000366) oligopeptide permease homolog AII [Borrelia burgdorferi]	290	3.90E-34
f6-21_aa	gi 2281455	(AF000365) oligopeptide permease homolog AIV [Borrelia burgdorferi]	279	9.90E-34
f6-21_aa	gi 2690261	(AE000790) oligopeptide ABC transporter, periplasmic	282	5.30E-33
f6-21_aa	gi 1616644	P30 [Borrelia burgdorferi]	271	6.70E-32
f6-21_aa	gi 2688226	(AE001139) oligopeptide ABC transporter, periplasmic	268	5.00E-31
f6-21_aa	gi 2281459	(AF000366) oligopeptide permease homolog AIII [Borrelia	268	5.00E-31
f6-21_aa	gi 2809546	(AF043071) oligopeptide permease periplasmic binding protein	268	5.00E-31
f6-21_aa	bbstl161785	60 kDa antigen [Borrelia coriaceae, C053, ATCC4338, Peptide, 514	255	2.90E-30
f6-21_aa	gi 2983834	(AE000740) transporter (extracellular solute binding protein family	154	3.50E-14
f6-27_aa	gi 2689911	(AE000792) B. burgdorferi predicted coding region BBB09 [Borrelia	1773	7.30E-240
f6-5_aa	gi 2689905	(AE000792) B. burgdorferi predicted coding region BBB27 [Borrelia	932	7.50E-126
f600_aa	gi 2281461	(AF000366) oligopeptide permease homolog C [Borrelia burgdorferi]	731	1.40E-100
f600_aa	gi 2688244	(AE001140) oligopeptide ABC transporter, permease protein (oppC-1)	731	1.40E-100
f600_aa	gi 143606	sporulation protein [Bacillus subtilis] >pirlC38447lC38447	372	5.00E-48
f600_aa	gi 40007	OppC gene product [Bacillus subtilis] >gnlPDIe1183165 oligopeptide	372	5.00E-48
f600_aa	gi 1574677	oligopeptide transport system permease protein (oppC)C [Haemophilus	372	7.30E-48
f600_aa	gi 47804	Opp C (AA1-301) [Salmonella typhimurium] >pirlC29333lQREBOC	366	4.20E-47
f600_aa	gnlPDIe10	Oligopeptide transport system permease protein OppC.	366	4.20E-47
f600_aa	15493	gnlPDIe11 (AJ002571) DppC [Bacillus subtilis] >gnlPDIe1183314	267	1.70E-42
f600_aa	81495			

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f600_aa	gi 1732315	transport system permease homolog [Listeria monocytogenes]	335	5.30E-42
f600_aa	gi 580851	dciAC [Bacillus subtilis] >sp P26904 DPPC_BACSU DIPEPTIDE TRANSPORT	258	1.50E-40
f600_aa	gn PPIdd10	oligopeptide transport system permease protein [Synechocystis 11164	240	2.50E-39
f600_aa	gi 677947	AppC [Bacillus subtilis] >gn PPIde1183160 oligopeptide ABC	236	2.80E-37
f600_aa	gi 1813497	di peptide transporter protein dppC [Bacillus firmus]	281	1.20E-35
f600_aa	sp Q10623	PUTATIVE PEPTIDE TRANSPORT PERMEASE PROTEIN	290	1.50E-35
f600_aa	Y021_MYC	CY373.01C.		
f600_aa	TU			
f600_aa	gi 1532201	BdKA [Streptomyces coelicolor]	291	1.60E-35
f603_aa	gi 2281460	(AF000366) oligopeptide permease homolog B [Borrelia burgdorferi]	1522	5.80E-214
f603_aa	gi 1574678	di peptide transport system permease protein (dppB) [Haemophilus	392	1.30E-100
f603_aa	gn PPIde11	oligopeptide ABC transporter (permease) [Bacillus subtilis]	374	3.40E-96
f603_aa	83164			
f603_aa	gi 580897	OppB gene product [Bacillus subtilis] >gi S15231 B38447	373	6.60E-96
f603_aa	gi 47803	Opp B (AA1-306) [Salmonella typhimurium] >gi B29333 QREBOB	371	6.70E-96
f603_aa	gi 1787497	(AE000223) oligopeptide transport system permease protein OppB	364	3.50E-95
f603_aa	gn PPIdd10	Oligopeptide transport system permease protein OppB.	357	3.50E-94
f603_aa	15492			
f603_aa	gi 580850	dcIA [Bacillus subtilis] >gn PPIde1181494 (AJ002571) DppB	350	9.10E-90
f603_aa	gi 551726	sporulation protein [Bacillus subtilis] >gi 143605 sporulation	374	2.40E-87
f603_aa	gi 349226	transmembrane protein [Escherichia coli] >gi 466682 dppB	293	9.60E-79
f603_aa	gi 1787053	(AE000185) o306; This 306 aa ORF is 46 pct identical (32 gaps) to	284	3.80E-77
f603_aa	gi 972895	DppB [Haemophilus influenzae] >gi 1574114 di peptide transport system	301	2.50E-76
f603_aa	gi 2182646	(AE000098) Y4tP [Rhizobium sp. NGR234] >sp Q53191 Y4TP_RHISN	294	9.10E-74
f603_aa	gi 2983140	(AE000692) transporter (OppBC family) [Aquifex aeolicus]	169	2.30E-73
f603_aa	gi 677946	AppB [Bacillus subtilis] >gn PPIde1183159 oligopeptide ABC	218	8.70E-73
f604_aa	gi 2281459	(AF000366) oligopeptide permease homolog AIII [Borrelia	2818	0
f604_aa	gi 2809546	(AF043071) oligopeptide permease periplasmic binding protein	2818	0
f604_aa	gi 2688226	(AE001139) oligopeptide ABC transporter, periplasmic	2823	0
f604_aa	gi 2688227	(AE001139) oligopeptide ABC transporter, periplasmic	1738	1.40E-234

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f604_aa	gi 2281458 [AF000366] oligopeptide permease homolog AII [Borrelia burgdorferi]	1731	1.30E-233
f604_aa	gi 2281468 [AF000366] OppAIV [Borrelia burgdorferi] >gi 2689891 (AE000792)	1675	3.60E-229
f604_aa	gi 2688228 [AE001139] oligopeptide ABC transporter, periplasmic	718	1.60E-204
f604_aa	gi 2809544 [AF043071] oligopeptide permease periplasmic binding protein	718	3.00E-204
f604_aa	gi 2253286 [AF005657] plasminogen binding protein [Borrelia burgdorferi]	718	4.10E-204
f604_aa	gi 2281457 [AF000366] oligopeptide permease homolog AI [Borrelia burgdorferi]	714	2.00E-203
f604_aa	bbsl 61785 60 kda antigen [Borrelia coriaceae, C053, ATCC 4338, Peptide, 514]	704	1.20E-190
f604_aa	gi 2281455 [AF000365] oligopeptide permease homolog AV [Borrelia burgdorferi]	1402	1.80E-188
f604_aa	gi 2690261 [AE000790] oligopeptide ABC transporter, periplasmic	1400	3.40E-188
f604_aa	gi 1616644 P30 [Borrelia burgdorferi]	858	4.90E-117
f604_aa	gi 47802 Opp A (AA1-542) [Salmonella typhimurium] >gi 47808 precursor	296	9.00E-114
f606_aa	gi 2281458 [AF000366] oligopeptide permease homolog AII [Borrelia burgdorferi]	2762	0
f606_aa	gi 2688227 [AE001139] oligopeptide ABC transporter, periplasmic	2774	0
f606_aa	gi 2281468 [AF000948] OppAIV [Borrelia burgdorferi] >gi 2689891 (AE000792)	1817	6.50E-245
f606_aa	gi 2809546 [AF043071] oligopeptide permease periplasmic binding protein	1739	3.10E-234
f606_aa	gi 2688226 [AE001139] oligopeptide ABC transporter, periplasmic	1738	4.20E-234
f606_aa	gi 2281459 [AF000366] oligopeptide permease homolog AIII [Borrelia	1733	2.00E-233
f606_aa	bbsl 61785 60 kda antigen [Borrelia coriaceae, C053, ATCC 4338, Peptide, 514]	762	1.70E-202
f606_aa	gi 2281455 [AF000365] oligopeptide permease homolog AV [Borrelia burgdorferi]	1456	1.80E-195
f606_aa	gi 2690261 [AE000790] oligopeptide ABC transporter, periplasmic	1454	3.30E-195
f606_aa	gi 2253286 [AF005657] plasminogen binding protein [Borrelia burgdorferi]	751	2.00E-192
f606_aa	gi 2688228 [AE001139] oligopeptide ABC transporter, periplasmic	751	2.70E-192
f606_aa	gi 2809544 [AF043071] oligopeptide permease periplasmic binding protein	751	6.90E-192
f606_aa	gi 2281457 [AF000366] oligopeptide permease homolog AI [Borrelia burgdorferi]	748	2.40E-191
f606_aa	gi 1616644 P30 [Borrelia burgdorferi]	1220	7.30E-163
f606_aa	gi 47802 Opp A (AA1-542) [Salmonella typhimurium] >gi 47808 precursor	285	7.80E-106
f607_aa	gi 2281457 [AF000366] oligopeptide permease homolog AI [Borrelia burgdorferi]	2694	0
f607_aa	gi 2253286 [AF005657] plasminogen binding protein [Borrelia burgdorferi]	2706	0
f607_aa	gi 2809544 [AF043071] oligopeptide permease periplasmic binding protein	2708	0
f607_aa	gi 2688228 [AE001139] oligopeptide ABC transporter, periplasmic	2714	0
f607_aa	bbsl 61785 60 kda antigen [Borrelia coriaceae, C053, ATCC 4338, Peptide, 514]	1272	3.80E-242

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f607_aa	gi 2809546	(AF043071) oligopeptide permease periplasmic binding protein	718	1.40E-204
f607_aa	gi 2688226	(AE001139) oligopeptide ABC transporter, periplasmic	718	3.60E-204
f607_aa	gi 2281459	(AF000366) oligopeptide permease homolog AIII [Borrelia	713	1.70E-203
f607_aa	gi 2688227	(AE001139) oligopeptide ABC transporter, periplasmic	751	2.40E-192
f607_aa	gi 2281458	(AF000366) oligopeptide permease homolog AII [Borrelia burgdorferi]	751	4.50E-192
f607_aa	gi 2281468	(AF000948) OppAIV [Borrelia burgdorferi] >gi 2689891 (AE000792)	806	8.40E-189
f607_aa	gi 2690261	(AE000790) oligopeptide ABC transporter, periplasmic	601	1.20E-144
f607_aa	gi 2281455	(AF000365) oligopeptide permease homolog AV [Borrelia burgdorferi]	600	1.60E-144
f607_aa	gi 1616644	P30 [Borrelia burgdorferi]	709	5.40E-103
f607_aa	gi 47802	Opp A (AA1-542) [Salmonella typhimurium] >gi 47808 precursor	261	8.50E-69
f611_aa	gi 2688231	(AE001139) B. burgdorferi predicted coding region BB0325 [Borrelia	1907	1.10E-261
f617_aa	gi 2688213	(AE001138) conserved hypothetical integral membrane protein	1574	2.70E-226
f617_aa	gi 2649711	(AE001042) ribose ABC transporter, permease protein (rbsC-1)	109	7.00E-12
f631_aa	gi 1165286	FtsW [Borrelia burgdorferi] >gi 2688164 (AE001137) cell division	1820	4.00E-259
f631_aa	gn IPDle22	membrane protein [Borrelia burgdorferi] >gn IPDle228289 ftsW	1815	2.10E-258
f631_aa	gi 146039	cell division protein [Escherichia coli] >gi 40857 FtsW protein	362	1.30E-60
f631_aa	gi 580938	internal open reading frame (AA 1-290) [Bacillus subtilis]	407	4.90E-55
f631_aa	gn IPDle31	FtsW [Mycobacterium tuberculosis] >sp O06223 FTW_H_MYCTU	412	5.40E-55
f631_aa	gi 580937	spoVE gene product (AA 1-366) [Bacillus subtilis] >gn IPDle1185111	410	2.90E-53
f631_aa	gi 143657	endospore forming protein [Bacillus subtilis]	405	1.20E-52
f631_aa	gn IPDle10	rod-shape-determining protein [Synechocystis sp.]	358	3.10E-51
f631_aa	gn IPDle12	(AL022602) cell division protein FtsW [Mycobacterium leprae]	396	6.70E-51
f631_aa	87793			
f631_aa	gi 1016213	strong sequence similarity to FtsW, RodA, and SpoV-E [Cyanophora	349	1.00E-50
f631_aa	gi 1574692	cell division protein (ftsW) [Haemophilus influenzae]	304	4.20E-50
f631_aa	85075	similar to cell-division protein [Bacillus subtilis]	281	1.80E-46
f631_aa	gi 1469784	putative cell division protein ftsW [Enterococcus hirae]	247	1.60E-38

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f631.aa	gil1572976	rod shape-determining protein (mreB) [Haemophilus influenzae]	196	1.20E-37
f631.aa	gil147695	rod-shape-determining protein [Escherichia coli] >gil1778551	194	5.00E-35
f635.aa	gil1165282	orf7; Method: conceptual translation supplied by author [Borrelia	1166	1.00E-156
f635.aa	gil1165282	ORF 224; The predicted gene product showed weak homology with the	621	2.80E-125
f647.aa	gil1448949	[Borrelia burgdorferi]	1032	1.00E-140
f647.aa	gil2688180	(AE001137) flagellar protein (flhB) [Borrelia burgdorferi]	1031	1.50E-140
f647.aa	gil1196323	putative [Borrelia burgdorferi]	1019	7.10E-139
f647.aa	gil1165270	orf19; Method: conceptual translation supplied by author [Borrelia	200	4.70E-24
f647.aa	gil2108242	22.5K protein [Treponema pallidum]	1095	8.10E-148
f65.aa	gil2688737	(AE001178) B. burgdorferi predicted coding region BB0792 [Borrelia	1220	1.70E-164
f653.aa	gil1165265	MotB [Borrelia burgdorferi] >gil185054 flagellar motor apparatus	168	5.80E-57
f653.aa	gil139286	MotB [Treponema phagedenis]	179	1.30E-49
f653.aa	gil2196896	MotB [Treponema pallidum]	1430	1.90E-199
f664.aa	gil1185062	flagellar export protein [Borrelia burgdorferi]	1430	1.90E-199
f664.aa	gil1165257	FlhB [Borrelia burgdorferi] >gil2688194 (AE001137) flagellar	1430	1.90E-199
f664.aa	gil1216382	FlhB' [Treponema pallidum] >pirPC4115[PC4115 flagellar protein	272	5.30E-64
f664.aa	gil395390	flagellar biosynthetic protein [Bacillus subtilis]	433	1.30E-61
f664.aa	gnlIPD1e11	flagella-associated protein [Bacillus subtilis]	433	1.30E-61
	85229		325	2.20E-39
f664.aa	gil1147737	third gene in flhQ operon; membrane protein homolog [Caulobacter	353	1.70E-46
f664.aa	gil2313898	(AE000589) flagellar biosynthetic protein (flhB) [Helicobacter	203	1.20E-44
f664.aa	gil2984250	(AE000768) flagellar biosynthetic protein FlhB [Aquitex aolicus]	319	3.00E-44
f664.aa	gil2459702	FlhB [Agrobacterium tumefaciens]	347	6.20E-44
f664.aa	gil793892	flhB [Yersinia enterocolitica] >pir[S54213 S54213 flhB protein -	330	1.30E-39
f664.aa	gnlIPD1d10	Flagellar biosynthetic protein FlhB. [Escherichia coli]		
	16420			
f664.aa	gil475126	YscU [Yersinia pseudotuberculosis] >gil2996233 (AF053946) Yop	309	9.80E-38
f664.aa	gil497216	YscU [Yersinia enterocolitica]	308	1.40E-37
f664.aa	gnlIPD1d10	flagellar protein FlhB [Salmonella typhimurium]	312	2.10E-37
f664.aa	07477			
f664.aa	gnlIPD1e28	secretion system apparatus, SsaU [Salmonella typhimurium]	312	8.20E-37
	3684			

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f679_aa	gil2688158	(AE001136) <i>B. burgdorferi</i> predicted coding region BB0259 [Borrelia	3714	0
f679_aa	gnlIPDd10	soluble lytic transglycosylase [Synechocystis sp.]	180	1.10E-25
f679_aa	gnlIPDd11	similar to lytic transglycosylase [Bacillus subtilis]	108	2.10E-22
	83177			
f679_aa	gil2984090	(AE000756) hypothetical protein [Aquifex aeolicus]	111	9.30E-17
f680_aa	gil2688153	(AE001136) bacitracin resistance protein (bacA) [Borrelia	769	3.90E-109
f680_aa	gnlIPDd11	similar to bacitracin resistance protein (undecaprenol	174	7.30E-18
	85988			
f680_aa	gil2622542	(AE000905) bacitracin resistance protein [Methanobacterium	116	3.30E-16
f680_aa	gil2984378	(AE000777) undecaprenol kinase [Aquifex aeolicus]	152	3.90E-15
f680_aa	gil882579	CG Site No. 29739 [Escherichia coli] >gil1789437 (AE000387)	139	2.60E-12
f688_aa	gil2688146	(AE001135) conserved hypothetical integral membrane protein	2497	0
f688_aa	gil2649351	(AE001019) conserved hypothetical protein [Archaeoglobus fulgidus]	110	3.70E-18
f688_aa	gil1592186	M. jannaschii predicted coding region MJ1562 [Methanococcus	174	1.10E-16
f7-30_aa	gil2690009	(AE000786) conserved hypothetical protein [Borrelia burgdorferi]	682	1.90E-90
f704_aa	gil2688137	(AE001134) glycerol uptake facilitator (glpF) [Borrelia	1307	4.70E-181
f704_aa	gil1429997	glycerol uptake facilitator [Bacillus subtilis] >gnlIPDd1182917	191	1.50E-50
f704_aa	gil521003	C01G6.1 [Caenorhabditis elegans]	152	1.60E-50
f704_aa	gil529582	water channel protein [Rattus norvegicus] >pir159266159266 water	142	5.80E-50
f704_aa	dbj1AB0005	(AB000507) aquaporin 7 [Rattus norvegicus]	155	1.30E-49
	07_1			
f704_aa	pirA57119	aquaporin 3 - human	149	4.20E-44
	A57119			
f704_aa	gil1109920	coded for by <i>C. elegans</i> cDNA cm16b11; strong similarity to MIP	168	9.30E-44
f704_aa	gnlIPDd10	aquaporin 3 [Homo sapiens] >sp Q92482 AQP3_HUMAN	148	5.30E-43
	19987			
f704_aa	gnlIPDd10	(AB008775) aquaporin 9 [Homo sapiens]	144	1.40E-42
	25786			
f704_aa	gil146188	glycerol diffusion facilitator [Escherichia coli] >gil305030 CG Site	146	1.30E-40
f704_aa	gil1065485	strong similarity to the MIP family of transmembrane channel	179	1.40E-39
f704_aa	sp P31140	GLYCEROL UPTAKE FACILITATOR PROTEIN	146	3.30E-39

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

GLPF_SHI FL			
f704.aa	gi 2587035 (AF026270) PduF [Salmonella typhimurium] >sp P37451 PDUF_SALTY	168	7.30E-39
f704.aa	gi 1399489 glycerol diffusion facilitator [Pseudomonas aeruginosa]	154	7.90E-39
f704.aa	gi 2649144 (AE001005) glycerol uptake facilitator, MFP channel (glpF)	150	1.30E-38
f707.aa	gi 2688143 (AE001134) B. burgdorferi predicted coding region BB0238 [Borrelia	1300	3.90E-176
f709.aa	gi 2688131 (AE001133) B. burgdorferi predicted coding region BB0236 [Borrelia	3437	0
f730.aa	gi 2688111 (AE001132) gufA protein [Borrelia burgdorferi] >pir C70127 C70127	1376	3.00E-192
f730.aa	gi 1707057 coded for by C. elegans cDNA CEESS55F; coded for by C. elegans cDNA	235	2.80E-83
f730.aa	gi 2621542 (AE000831) conserved protein [Methanobacterium thermoautotrophicum]	259	1.10E-74
f730.aa	gn P Dle18 gufA gene product [Myxococcus xanthus] >gi 49253 orfX gene 3440	175	2.30E-35
f730.aa	gi 2984109 (AE000757) hypothetical protein [Aequifex aeolicus]	171	7.00E-28
f736.aa	gi 2688115 (AE001132) phosphate ABC transporter, periplasmic phosphate-binding	1403	2.10E-186
f736.aa	gi 2622858 (AE000929) phosphate-binding protein PstS [Methanobacterium	151	4.40E-30
f736.aa	gi 2622859 (AE000929) phosphate-binding protein PstS homolog [Methanobacterium	145	2.80E-24
f736.aa	gn P Dd10 ORF108 [Bacillus subtilis] >gn P D e1185766 alternate gene 10224	120	1.20E-11
f739.aa	gi 2688119 (AE001132) B. burgdorferi predicted coding region BB0213 [Borrelia	1139	1.10E-156
f742.aa	gi 2688100 (AE001131) surface-located membrane protein 1 (lmp1) [Borrelia	5654	0
f742.aa	gi 2621120 (AE000799) O-linked GicNAc transferase [Methanobacterium	200	9.30E-22
f742.aa	gi 2621106 (AE000798) O-linked GicNAc transferase [Methanobacterium	180	5.80E-17
f742.aa	pir E69190I conserved hypothetical protein MTH68 - Methanobacterium E69190	154	1.60E-14
f742.aa	gi 1591608 transformation sensitive protein [Methanococcus jannaschii]	109	9.90E-14
f742.aa	gi 1589778 SPINDLY (Arabidopsis thaliana)	101	1.40E-13
f742.aa	gi 2984175 (AE000762) hypothetical protein [Aequifex aeolicus]	132	7.30E-13
f742.aa	gi 3037137 (AF056198) Hsp70/Hsp90 organizing protein homolog [Drosophila	105	5.40E-11
f743.aa	gi 2688104 (AE001131) B. burgdorferi predicted coding region BB0209 [Borrelia	1299	1.70E-174
f748.aa	gi 2688089 (AE001130) Lambda CII stability-governing protein (hflC) [Borrelia	1615	5.10E-220

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f748.aa	gi 436158	putative integral membrane protease required for high frequency	191	4.80E-35
f748.aa	gi 1573107	Lambda CII stability-governing protein (hfcC) [Haemophilus	193	4.90E-33
f748.aa	gi 507735	HfcC [Vibrio parahaemolyticus] >sp P40606 HFLC_VIBPA_HFLC	212	6.10E-26
f752.aa	gi 2688092	(AE001130)	2585	0
f752.aa	gi 2984050	((AE000754)) UDP-MurNAc-tripeptide synthetase [Aquifex aeolicus]	202	9.10E-74
f752.aa	gi 40162	murE gene product [Bacillus subtilis] >gi 1185108	157	6.40E-70
f752.aa	gi 1PD1d10	UDP-MurNAc-tripeptide synthetase [Synechocystis sp.]	166	5.20E-57
f752.aa	gi 1PD1d30	UDP-MurNAc-tripeptide synthetase [Rickettsia prowazekii]	108	2.30E-51
f752.aa	gi 1574688	UDP-MurNAc-tripeptide synthetase (mure) [Haemophilus influenzae]	166	3.20E-50
f752.aa	gi 1PD1d12	(AL022602) udp-n-acetyl muramoylalanyl-d-glutamate	183	3.20E-50
f752.aa	gi 1PD1d31	MurE [Mycobacterium tuberculosis]	181	4.10E-46
f752.aa	gi 581032	UDP-MurNAc-tripeptide synthetase (MurE) [Escherichia coli]	175	1.30E-41
f752.aa	gi 2177098	UDP-MurNAc-Dipeptid: meso-diaminopimelate ligase [Escherichia	172	3.70E-41
f752.aa	gi 2314673	((AE000648)) UDP-MurNAc-tripeptide synthetase (murE) [Helicobacter	137	9.80E-41
f752.aa	gi 840843	UDP-N-acetyl muramoylalanyl-D-glutamate--2,6-diaminopimelate ligase	135	1.70E-20
f76-1.aa	gi 1209837	lipoprotein [Borrelia burgdorferi]	395	2.80E-49
f76-1.aa	gi 1209873	lipoprotein [Borrelia burgdorferi]	250	7.00E-37
f76-1.aa	gi 1209843	lipoprotein [Borrelia burgdorferi]	267	7.30E-32
f76-1.aa	gi 21280	(AF000270) lipoprotein [Borrelia burgdorferi] >gi 3095109	258	1.20E-30
f76-1.aa	gi 1PD1d26	surface-exposed lipoprotein [Borrelia afzelii]	116	2.40E-18
f76-1.aa	gi 1209849	lipoprotein [Borrelia burgdorferi]	146	8.30E-17
f76-1.aa	gi 3095105	(AF046998) 2.9-8 lipoprotein [Borrelia burgdorferi]	148	5.80E-14
f76-1.aa	gi 3095107	(AF046999) 2.9-9 lipoprotein [Borrelia burgdorferi]	127	7.20E-11
f764.aa	gi 2688084	(AE001129) B. burgdorferi predicted coding region BB0193 [Borrelia	1218	1.20E-164
f770.aa	gi 2688077	(AE001129) conserved hypothetical protein [Borrelia burgdorferi]	646	7.60E-87
f790.aa	gi 2688065	(AE001128) outer membrane protein (tpn50) [Borrelia burgdorferi]	2013	2.50E-271

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f790_aa	gi 458015	TPN50 precursor [Treponema pallidum]		134	4.30E-33
f790_aa	sp P38369 T_P50_TREP_A	OUTER MEMBRANE PROTEIN TPN50 PRECURSOR.		134	4.30E-33
f790_aa	gi 532658	antigen [Treponema pallidum]>pir S61867 S61867 antigen lpp57 -		139	4.30E-31
f792_aa	gi 2688052	(AE001127) B. burgdorferi predicted coding region BB0165 [Borrelia	3185	0	
f797_aa	gi 2688056	(AE001127) B. burgdorferi predicted coding region BB0159 [Borrelia	1116	5.30E-148	
f798_aa	gi 2688051	(AE001127) antigen, S2, putative [Borrelia burgdorferi]	1223	9.70E-164	
f798_aa	gi 1063419	S2 gene product [Borrelia burgdorferi]	116	4.70E-23	
f798_aa	gi 2690227	(AE000790) antigen, S2 [Borrelia burgdorferi] >pir D70207 D70207	116	1.50E-22	
f798_aa	gi 2690128	(AE000788) protein p23 [Borrelia burgdorferi] >pir C70257 C70257	110	1.40E-19	
f798_aa	gi 2689956	(AE000785) protein p23 [Borrelia burgdorferi] >pir D70225 D70225	104	2.70E-15	
f799_aa	gi 2688043	(AE001126) B. burgdorferi predicted coding region BB0156 [Borrelia	632	1.40E-83	
f8-10_aa	gi 2690052	(AE000784) antigen, P35, putative [Borrelia burgdorferi]	1241	1.10E-167	
f8-10_aa	gi 2689955	(AE000785) antigen, P35, putative [Borrelia burgdorferi]	298	1.70E-57	
f8-10_aa	gi 2690120	(AE000789) B. burgdorferi predicted coding region BB134 [Borrelia	254	3.80E-54	
f8-10_aa	gi 2690100	(AE000789) B. burgdorferi predicted coding region BB116 [Borrelia	182	2.90E-31	
f8-10_aa	gi 2690207	(AE000787) B. burgdorferi predicted coding region BB102 [Borrelia	196	1.50E-20	
f8-10_aa	gi 2690116	(AE000789) B. burgdorferi predicted coding region BB129 [Borrelia	192	5.50E-20	
f8-10_aa	gi 2690125	(AE000788) antigen, P35, putative [Borrelia burgdorferi]	129	5.80E-14	
f8-10_aa	gi 2690206	(AE000787) B. burgdorferi predicted coding region BB101 [Borrelia	103	1.10E-13	
f8-10_aa	gi 2690099	(AE000789) B. burgdorferi predicted coding region BB115 [Borrelia	142	8.50E-13	
f8-10_aa	gi 2690115	(AE000789) B. burgdorferi predicted coding region BB128 [Borrelia	130	3.30E-12	
f8-14_aa	gi 2690074	(AE000784) B. burgdorferi predicted coding region BBH37 [Borrelia	1560	2.60E-206	
f8-14_aa	gi 2690188	(AE000787) B. burgdorferi predicted coding region BB108 [Borrelia	599	3.50E-123	
f8-14_aa	gi 2690030	(AE000786) B. burgdorferi predicted coding region BBG01 [Borrelia	337	4.40E-106	
f8-14_aa	gi 2690139	(AE000788) B. burgdorferi predicted coding region BBK01 [Borrelia	173	8.00E-91	
f8_aa	gi 2688783	(AE001182) B. burgdorferi predicted coding region BB0840 [Borrelia	2765	0	
f8_aa	gi 2697112	(AF008219) unknown [Borrelia afzelii]	1494	2.80E-205	
f800_aa	gi 2688044	(AE001126) B. burgdorferi predicted coding region BB0155 [Borrelia	1936	1.00E-262	
f805_aa	gi 2688039	(AE001126) N-acetylglucosamine-6-phosphate acetylase (nagA)	641	6.30E-85	

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f810_aa	gi 2688024	(AE001125) glycine betaine, L-proline ABC transporter,		1527	4.20E-207
f810_aa	gi 984805	glycine betaine-binding protein precursor [Bacillus subtilis]		179	6.80E-21
f810_aa	gi 1850605	ProX [Streptococcus mutans]		181	2.30E-18
f814_aa	pir D701171	acriflavine resistance protein (acrB) homolog - Lyme disease D70117	5105	0	
f814_aa	gi 2688027	(AE001125) acriflavine resistance protein (acrB) [Borrelia	5111	0	
f814_aa	gi 2983346	(AE000707) cation efflux (AcrB/AcrD/AcrF family) [Aquifex aeolicus]	325	4.80E-119	
f814_aa	gi 2313726	(AE000574) acriflavine resistance protein (acrB) [Helicobacter	327	4.50E-111	
f814_aa	gi 3068786	(AF059041) RND pump protein [Helicobacter pylori]	297	1.70E-110	
f814_aa	gnl P D e11	similar to acriflavine resistance protein [Bacillus subtilis] 82651	257	8.90E-100	
f814_aa	gi 1573914	acriflavine resistance protein (acrB) [Haemophilus influenzae]	294	2.10E-97	
f814_aa	gnl P D e25	mexF [Pseudomonas aeruginosa] 6815	300	2.00E-88	
f814_aa	gnl P D d10	cation efflux system protein CzcA [Synechocystis sp.] 19295	198	1.30E-87	
f814_aa	gnl P D e28	membrane-bound cation-proton-antiporter [Ralstonia eutropha] 5274	283	2.20E-87	
f814_aa	gi 438854	envD homologue; ORFB [Pseudomonas aeruginosa] >pir S39630 S39630	290	6.50E-87	
f814_aa	gnl P D d10	CzcA [Alcaligenes sp.] >pir JC4700 JC4700 cadmium, zinc, 11721	275	8.20E-87	
f814_aa	gi 2314107	(AE000605) cation efflux system protein (czcA) [Helicobacter	266	2.30E-86	
f814_aa	pir A338301	cation efflux system membrane protein czcA - Alcaligenes A33830	275	3.10E-86	
f814_aa	gnl P D d10	envD gene product homolog [Escherichia coli] >gi 1788814 17073	283	8.30E-86	
f818_aa	gi 2688032	(AE001125) B. burgdorferi predicted coding region BB0139 [Borrelia	664	3.00E-87	
f82_aa	gi 2688729	(AE001177) B. burgdorferi predicted coding region BB0776 [Borrelia	991	2.20E-132	
f820_aa	gi 2688029	(AE001125) penicillin-binding protein (ppb-1) [Borrelia	3171	0	
f820_aa	gi 580936	SpoVD [Bacillus subtilis] >gnl P D e1185107 penicillin-binding	149	3.00E-49	
f820_aa	gi 150283	penicillin-binding protein 2 [Neisseria meningitidis]	154	6.90E-43	
f820_aa	gnl P D e12	(AL022602) penicillin binding protein 2 [Mycobacterium	182	4.20E-42	

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

	87798					
f820_aa	gi 509190	penicillin-binding protein 2 [Neisseria meningitidis]			158	1.70E-41
f820_aa	gi 509118	penicillin-binding protein 2 [Neisseria meningitidis]			151	7.10E-41
f820_aa	gi 840842	penicillin-binding protein 3 [Pseudomonas aeruginosa]			177	1.20E-40
f820_aa	gi 509065	penicillin-binding protein 2 [Neisseria meningitidis]			152	1.40E-40
f820_aa	gi 509043	penicillin-binding protein 2 [Neisseria meningitidis]			150	2.70E-40
f820_aa	gi 509159	penicillin-binding protein 2 [Neisseria meningitidis]			147	2.80E-40
f820_aa	gi 509120	penicillin-binding protein 2 [Neisseria meningitidis]			155	1.60E-39
f820_aa	gi 509157	penicillin-binding protein 2 [Neisseria meningitidis]			155	1.60E-39
f820_aa	gi 509126	penicillin-binding protein 2 [Neisseria meningitidis]			158	1.70E-39
f820_aa	gi 45178	penicillin-binding protein 2 (AA 1 - 581) [Neisseria meningitidis]			155	2.30E-38
f820_aa	gi 150279	penicillin binding protein 2 [Neisseria gonorrhoeae]			154	8.70E-38
f831_aa	gi 2688018	(AE001124) B. burgdorferi predicted coding region BB0126 [Borrelia			994	1.20E-133
f843_aa	gi 2688014	(AE001124) PTS system, maltose and glucose-specific IIABC component	2590	0		
f843_aa	gi 2688579	(AE001166) PTS system, glucose-specific IIBC component (ptsG)			594	1.80E-129
f843_aa	gi 1072418	glcA [Staphylococcus carnosus] >pirS46952 S46952			283	1.00E-72
f843_aa	gi 1072419	glcB [Staphylococcus carnosus] >pirS63606 S46953			248	1.00E-66
f843_aa	dbj ID86417	Yfif [Bacillus subtilis] >gnl P D e1182760 similar to			215	7.90E-65
	1					
f843_aa	gi 2197104	(AF003742) MalX homolog [Escherichia coli]			182	8.90E-64
f843_aa	gi 43819	nagE gene product [Klebsiella pneumoniae] >pirS18607 S18607			264	8.50E-63
f843_aa	gi 146913	N-acetyl glucosamine transport protein [Escherichia coli]			256	1.10E-62
f843_aa	gi 39956	IIGlc [Bacillus subtilis] >gnl P D e11849797 phosphotransferase system			315	5.20E-62
f843_aa	dbj ID87820	NagE [Vibrio cholerae non-O1] >pir C5651 C5651			263	3.80E-61
	1					
f843_aa	gi 2689888	(AE000792) PTS system, maltose and glucose-specific IIABC component			198	1.10E-60
f843_aa	gi 397363	enzyme II-glc [Salmonella typhimurium] >pirS36620 S36620			227	1.20E-58
f843_aa	gi 147393	glucose-specific enzyme II of phosphotransferase system [Escherichia			226	3.90E-57
f843_aa	gnl P D e1182187	alternate gene name: yzfA; similar to phosphotransferase			180	9.00E-56
f843_aa	gi 1732194	PTS permease for glucose [Vibrio furnissii]			349	4.30E-50

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f850.aa	gi 2687999 (AE001123) <i>B. burgdorferi</i> predicted coding region BB0110 [Borrelia]	2374	0
f853.aa	gi 2687994 (AE001123) basic membrane protein [Borrelia burgdorferi]	1672	2.20E-224
f853.aa	gi 155055 basic membrane protein precursor [Treponema pallidum]	130	3.60E-24
f859.aa	gi 2688002 (AE001123) <i>B. burgdorferi</i> predicted coding region BB0102 [Borrelia]	888	1.80E-115
f86.aa	gi 2688725 (AE001177) flagellar P-ring protein (flgD) [Borrelia burgdorferi]	1647	1.50E-217
f86.aa	gi 2920802 (AF019213) FlgI [Vibrio cholerae]	143	3.50E-14
f86.aa	gi 405550 flagellar P-ring protein [Pseudomonas putida] >sp Q52082 FLGL_PSEPU	102	3.70E-13
f86.aa	gi 144241 flagellin [Caulobacter crescentus] >pir A41891 A41891 basal body	110	6.70E-13
f860.aa	gi 2687998 (AE001123) asparaginyl-tRNA synthetase (asnS) [Borrelia]	1110	2.40E-149
f860.aa	gi 1574761 asparaginyl-tRNA synthetase (asnS) [Haemophilus influenzae]	634	1.30E-83
f860.aa	gi 147935 asparaginyl-tRNA synthetase (asnS) [Escherichia coli] >gi 41000	622	6.10E-82
f860.aa	gn IPD 12 (AJ222644) asparaginyl-tRNA synthetase [Arabidopsis thaliana] 02698	404	2.40E-80
f860.aa	gn IPD 10 asparaginyl-tRNA synthetase [Synechocystis sp.] 11495	618	4.50E-80
f860.aa	gi 530408 Asn-tRNA synthetase [Mycoplasma capricolum] >pir S77842 S77842	439	1.60E-65
f860.aa	gi 1045792 asparaginyl-tRNA synthetase [Mycoplasma genitalium]	365	2.20E-62
f860.aa	gi 1674281 (AE000057) Mycoplasma pneumoniae, asparaginyl-tRNA synthetase;	338	3.10E-61
f860.aa	gn IPD 12 (AJ222645) asparaginyl-tRNA synthetase [Arabidopsis thaliana] 02700	364	3.90E-59
f860.aa	gn IPD 26 YCR024c, len:492 [Saccharomyces cerevisiae] >pir S19435 S19435	150	3.90E-47
f860.aa	gn IPD 25 asparaginyl-tRNA synthetase [Salmonella typhi] 4488	370	1.70E-46
f860.aa	gn IPD 4305 asparaginyl-tRNA synthetase [Salmonella typhi] 4305	224	1.30E-44
f860.aa	gn IPD 18 asparagine-tRNA ligase [Lactobacillus delbrueckii] 8505	224	1.30E-44
f860.aa	pir S71072 asparagine-tRNA ligase (EC 6.1.1.22) asnS1 - Lactobacillus S71072	224	1.30E-44
f860.aa	gn IPD 18 asparagine-tRNA ligase [Lactobacillus delbrueckii] 8572	224	2.40E-44
f860.aa	gi 1146247 asparaginyl-tRNA synthetase [Bacillus subtilis] >gn IPD 1183681	234	6.10E-44
f861.aa	gi 2687975 (AE001122) glutamate racemase (mru) [Borrelia burgdorferi]	1354	2.90E-186

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f861_aa	gii396314	glutamate synthase [Escherichia coli] >gii290428 glutamate synthase	168	1.20E-16
f861_aa	gnlIPD1e11	glutamate racemase [Bacillus subtilis] >gnlIPD1e1184088	120	1.80E-13
	65353			
f861_aa	pirJCS5871J	glutamate racemase (EC 5.1.1.3) - Bacillus pumilus	122	1.80E-13
	C5587			
f861_aa	spIP529731	PROBABLE GLUTAMATE RACEMASE (EC 5.1.1.3).	114	8.10E-13
	MURI_HA			
	EIN			
f867_aa	gii2687979	(AE001122) V-type ATPase, subunit A (atpA) [Borrelia burgdorferi]	2826	0
	pirJCS5321J	vacuolar-type ATPase (EC 3.7.7.1) A chain - Desulfurococcus	594	2.20E-162
	C5532			
f867_aa	gii2104726	V-ATPase A subunit [Desulfurococcus sp. SY]	594	3.10E-162
	gii2605627	ATPase alpha subunit [Thermococcus sp.]	592	7.10E-161
f867_aa	gnlIPD1d10	Na+ -ATPase alpha subunit [Enterococcus hirae]	601	1.60E-153
	03475			
f867_aa	gii1590955	H+-transporting ATP synthase, subunit A (atpA) [Methanococcus	585	6.00E-147
	gii496904	membrane ATPase [Haloferax volcanii] >pir S55895 S45144	728	6.00E-147
f867_aa	gii152927	ATPase alpha subunit [Sulfolobus acidocaldarius] >pir A28652 A28652	548	5.00E-163
	gii2649416	(AE001023) H+-transporting ATP synthase, subunit A (atpA)	748	2.00E-146
f867_aa	gii2622052	(AE000869) ATP synthase, subunit A [Methanobacterium	607	9.40E-146
f867_aa	gii168926	vacuolar ATPase vma-1 [Neurospora crassa] >pir A30799 PXNCV7	302	9.00E-145
	gii149820	ATPase alpha subunit [Methanoscarcina barkeri] >pir A34283 A34283	743	1.40E-143
f867_aa	gii160736	vacuolar ATPase [Plasmodium falciparum] >pir A48582 A48582 vacuolar	305	9.40E-140
f867_aa	gnlIPD1d10	adenosine triphosphatase A subunit [Acetabularia acetabulum]	307	9.00E-137
	09732			
f867_aa	gii49048	ATPase alpha-subunit [Thermus aquaticus thermophilus]	684	4.80E-136
	gii2687980	(AE001122) V-type ATPase, subunit B (atpB) [Borrelia burgdorferi]	2205	1.80E-298
f868_aa	gii1590954	H+-transporting ATP synthase, subunit B (atpB) [Methanococcus	156	2.00E-114
	gii2605628	ATPase beta subunit [Thermococcus sp.]	151	3.30E-108
f868_aa	gii2104727	V-ATPase B subunit [Desulfurococcus sp. SY]	151	1.10E-107
f868_aa	gii43641	ATP synthase subunit [Halobacterium salinarium] >pir S14733 S14733	150	1.80E-107
	gii149821	ATPase beta subunit [Methanoscarcina barkeri] >pir B34283 B34283	172	1.00E-105

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f868.aa	gnl P D d10 Na+ -ATPase beta subunit [Enterococcus hirae] 03476		151	1.40E-105
f868.aa	gi 2649415 (AE001023) H+-transporting ATP synthase, subunit B (atpB)		151	1.70E-103
f868.aa	gi 496905 membrane ATPase [Haloferax volcanii] >pir S55896 S45145		153	5.80E-103
f868.aa	gi 1199639 A1AO H+-ATPase, subunit B [Methanosarcina mazeii]		173	2.20E-102
f868.aa	gi 2622051 (AE000869) ATP synthase, subunit B [Methanobacterium		155	1.00E-101
f868.aa	gnl P D d10 adenosine triphosphatase B subunit [Acetabularia acetabulum] 09734		159	1.30E-101
f868.aa	gi 1086645 Similar to vacuolar ATP synthase (strong). [Caenorhabditis elegans]		163	1.30E-101
f868.aa	gi 459198 vacuolar H+-ATPase subunit B [Gossypium hirsutum]		164	4.60E-101
f868.aa	gi 167108 vacuolar ATPase B subunit [Hordeum vulgare] >sp Q40078 VAT1_HORVU		164	4.60E-101
f872.aa	gi 2687986 (AE001122) B. burgdorferi predicted coding region BB0089 [Borrelia		1684	1.60E-230
f874.aa	gi 2687965 (AE001121) L-lactate dehydrogenase (ldh) [Borrelia burgdorferi]		1603	2.80E-217
f874.aa	gi 39758 L-lactate dehydrogenase [Bacillus psychrosaccharolyticus]		520	3.10E-109
f874.aa	pir S08183 L-lactate dehydrogenase (EC 1.1.1.27) X - Bacillus S08183		515	4.30E-109
f874.aa	pir A25805 L-lactate dehydrogenase (EC 1.1.1.27) - Bacillus subtilis A25805		520	1.00E-107
f874.aa	gi 143136 L-lactate dehydrogenase [Bacillus megaterium] >pir S00133 DEBSLM		430	5.20E-107
f874.aa	gi 43138 lactate dehydrogenase (EC 1.1.1.27) [Bacillus stearothermophilus]		514	6.60E-107
f874.aa	gnl P D d10 L-lactate dehydrogenase [Bacillus subtilis] >gnl P D e 1182257 09574		512	8.90E-107
f874.aa	gi 143134 lactate dehydrogenase (EC 1.1.1.27) [Bacillus caldotenax]		516	1.70E-106
f874.aa	gi 143132 lactate dehydrogenase (AC 1.1.1.27) [Bacillus caldotenax]		506	2.30E-106
f874.aa	gi 412392 NAD-dependent dehydrogenase [unidentified]		508	4.40E-106
f874.aa	gi 143130 L-lactate dehydrogenase [Bacillus caldotenax] >pir S00019 S00019		510	1.10E-105
f874.aa	gi 642256 L-lactate dehydrogenase [Pediococcus acidilactici]		560	1.70E-91
f874.aa	gi 847956 L-lactate dehydrogenase [Lactobacillus sake] >sp P50934 LDH_LACSK		381	2.30E-91
f874.aa	gi 581305 L-lactate dehydrogenase [Lactobacillus plantarum] >pir A36957 A36957		547	2.30E-91
f874.aa	gi 149575 L(+)-lactate dehydrogenase [Lactobacillus casei]		386	3.20E-91
f886.aa	gi 2687958 (AE001120) B. burgdorferi predicted coding region BB0077 [Borrelia		1792	9.50E-237

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

Query	GenSeq Access No.	GenSeq Gene Description	BLAST Score	BLAST P-Value
f888.aa	gi 2687959 (AE001120) <i>B. burgdorferi</i> predicted coding region BB0075 [Borrelia	2351 3.599999944 710933e-318		
f893.aa	gi 2687962 (AE001120) <i>B. burgdorferi</i> predicted coding region BB0071 [Borrelia	2514 0		
f895.aa	gi 2687954 (AE001120) conserved hypothetical protein [Borrelia burgdorferi]	747 3.60E-100		
f895.aa	gnl PTD e11 similar to hypothetical proteins [Bacillus subtilis] 84285	103 2.50E-35		
f899.aa	gi 2687946 (AE001119) <i>B. burgdorferi</i> predicted coding region BB0066 [Borrelia	1161 4.30E-158		
f924.aa	gi 2687934 (AE001118) <i>B. burgdorferi</i> predicted coding region BB0044 [Borrelia	692 3.90E-93		
f925.aa	gi 2687935 (AE001118) <i>B. burgdorferi</i> predicted coding region BB0043 [Borrelia	1771 7.50E-242		
f929.aa	gi 2687916 (AE001117) <i>B. burgdorferi</i> predicted coding region BB0038 [Borrelia	2589 0		
f93.aa	gi 2688703 (AE001176) pyridoxal kinase (pdXK) [Borrelia burgdorferi]	1334 6.60E-181		
f933.aa	gi 2687917 (AE001117) <i>B. burgdorferi</i> predicted coding region BB0034 [Borrelia	902 1.90E-122		
f933.aa	gi 2690091 (AE000789) conserved hypothetical protein [Borrelia burgdorferi]	136 3.10E-37		
f933.aa	gi 2690225 (AE000790) conserved hypothetical protein [Borrelia burgdorferi]	149 4.50E-37		
f933.aa	gi 2690045 (AE000784) conserved hypothetical protein [Borrelia burgdorferi]	126 5.70E-28		
f933.aa	gi 22239281 No definition line found [Borrelia burgdorferi]	148 2.40E-14		
f939.aa	gi 2687919 (AE001117) <i>B. burgdorferi</i> predicted coding region BB0028 [Borrelia	1796 7.50E-241		
f940.aa	gi 2687920 (AE001117) <i>B. burgdorferi</i> predicted coding region BB0027 [Borrelia	1109 1.20E-152		
f943.aa	gi 2687905 (AE001116) <i>B. burgdorferi</i> predicted coding region BB0024 [Borrelia	2001 5.00E-273		
f943.aa	gi 411592 L-sorbose dehydrogenase [unidentified]	175 2.30E-15		
f943.aa	gnl PTD d10 L-sorbose dehydrogenase [Acetobacter liquefaciens] 06418	173 4.40E-15		
f952.aa	gi 2687880 (AE001115) glpE protein (glpE) [Borrelia burgdorferi]	628 2.90E-84		

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f147.aa	W06425	Water-forming NADH oxidase.	369	8.00E-86
f147.aa	R32089	Benzene dioxygenase polypeptide V.	104	4.70E-11
f147.aa	R66733	Aromatic dihydrodiol/catechol deoxygenase #5.	105	9.00E-11
f152.aa	R81549	High affinity potassium uptake transporter HKT1.	137	3.70E-18
f157.aa	W15192	Staphylococcus aureus cell surface protein.	239	3.40E-37
f17-6.aa	W30763	Mannose-1-phosphate transferase protein MNNA4.	178	5.20E-16
f17-6.aa	W03627	Human follicle stimulating hormone GPR N-terminal sequence.	145	1.30E-11
f17-6.aa	W03626	Human thyrotropin GPR N-terminal sequence.	144	1.90E-11
f17-6.aa	W21591	Antibiotic potentiating peptide #3.	141	5.10E-11
f196.aa	W05196	Helicobacter pylori 50 kDa protective antigen G3.8.	183	2.70E-18
f196.aa	W20916	H. pylori inner membrane protein 14gp12015orf12.	180	3.60E-17
f196.aa	W20287	H. pylori inner membrane protein, 24132293.aa.	169	6.50E-15
f196.aa	W20769	H. pylori inner membrane protein, 07ee20513orf28.	169	1.40E-14
f196.aa	W20767	H. pylori cytoplasmic protein, 07ee20513orf1.	140	6.10E-14
f197.aa	W20769	H. pylori inner membrane protein, 07ee20513orf28.	190	2.30E-19
f197.aa	W20287	H. pylori inner membrane protein, 24132293.aa.	190	2.00E-18
f197.aa	W05196	Helicobacter pylori 50 kDa protective antigen G3.8.	179	4.00E-16
f197.aa	W20916	H. pylori inner membrane protein 14gp12015orf12.	182	6.30E-16
f197.aa	W20767	H. pylori cytoplasmic protein, 07ee20513orf1.	150	1.10E-12
f21-4.aa	R69629	B. burgdorferi OspF operon.	321	7.00E-39
f21-4.aa	R89476	B. burgdorferi OspG lipoprotein.	107	6.10E-34
f24-1.aa	W22676	Borrelia variable major protein (VMP)-like protein VlsE.	412	4.60E-72
f291.aa	W20152	H. pylori transporter protein, 1464715.aa.	336	1.70E-41
f291.aa	W24682	Helicobacter pylori transporter protein 4882763.aa.	234	8.20E-27
f291.aa	W20528	H. pylori cell envelope transporter protein 4882763.aa.	234	8.20E-27
f291.aa	W20592	H. pylori transporter protein, 01ce11513orf21.	168	7.60E-17
f301.aa	W20287	H. pylori inner membrane protein, 24132293.aa.	158	1.60E-13
f301.aa	W20916	H. pylori inner membrane protein 14gp12015orf12.	158	1.90E-13
f301.aa	W20769	H. pylori inner membrane protein, 07ee20513orf28.	158	2.40E-13
f301.aa	W05196	Helicobacter pylori 50 kDa protective antigen G3.8.	157	2.80E-13
f301.aa	W20767	H. pylori cytoplasmic protein, 07ee20513orf1.	138	4.30E-11

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f320.aa	R24300.	Glycopeptide resistance protein VanY from <i>E.faecium</i> .		142	2.90E-14
f328.aa	R15642	CRP synthetase.	274	3.00E-50	
f328.aa	W20778	H. pylori cytoplasmic protein, 07ge2041Sorf6.	122	1.90E-34	
f352.aa	W03626	Human thyrotropin GPR N-terminal sequence.	153	4.70E-12	
f352.aa	W21591	Antibiotic potentiating peptide #3.	152	6.60E-12	
f352.aa	W03627	Human follicle stimulating hormone GPR N-terminal sequence.	145	5.30E-11	
f4-50.aa	W07187	B. garinii TP90 decorin binding protein.	305	1.30E-41	
f4-50.aa	W07186	B. afzelii strain pGau decorin binding protein.	161	1.60E-34	
f4-50.aa	W07185	B. burgdorferi HB-19 decorin binding protein.	173	2.80E-34	
f4-50.aa	W07183	B. burgdorferi B31 decorin binding protein.	176	1.80E-33	
f4-50.aa	W07190	B. burgdorferi JD1 decorin binding protein.	177	1.80E-33	
f4-50.aa	W07182	B. burgdorferi 297 decorin binding protein.	177	1.10E-32	
f4-50.aa	W07189	B. burgdorferi LP7 decorin binding protein.	177	1.10E-32	
f4-50.aa	W07188	B. burgdorferi LP4 decorin binding protein.	177	3.90E-32	
f4-50.aa	W07184	B. burgdorferi Sh.2.822 decorin binding protein.	177	1.30E-31	
f45-2.aa	R89476	B. burgdorferi OspG lipoprotein.	213	1.30E-35	
f45-2.aa	R70491	Leucocytozoan protozoa structural protein epitope.	206	2.10E-20	
f45-2.aa	W03626	Human thyrotropin GPR N-terminal sequence.	211	6.10E-20	
f45-2.aa	W03627	Human follicle stimulating hormone GPR N-terminal sequence.	202	8.90E-19	
f45-2.aa	R69629	B. burgdorferi OspF operon.	111	1.10E-14	
f45-2.aa	W30763	Mannose-1-phosphate transferase protein MNMN4.	166	1.00E-13	
f45-2.aa	R97866	Chicken leucocytozoan immunogenic protein for use in vaccines.	154	7.10E-12	
f488.aa	W15078	<i>M. leprae</i> gyra precursor.	390	2.70E-143	
f488.aa	R88733	<i>S. aureus</i> mutant grlA protein.	698	6.70E-122	
f488.aa	R88731	<i>S. aureus</i> topoisomerase IV grlA subunit.	698	6.70E-122	
f49-2.aa	W22676	Borrelia variable major protein (VMP)-like protein VlsE.	497	2.70E-75	
f5-14.aa	W03626	Human thyrotropin GPR N-terminal sequence.	234	6.60E-23	
f5-14.aa	W03627	Human follicle stimulating hormone GPR N-terminal sequence.	231	1.40E-22	
f5-14.aa	R70491	Leucocytozoan protozoa structural protein epitope.	221	1.00E-20	
f5-14.aa	W30763	Mannose-1-phosphate transferase protein MNMN4.	203	1.60E-18	
f5-14.aa	R97866	Chicken leucocytozoan immunogenic protein for use in vaccines.	187	2.10E-15	

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f5-14.aa	W21591	Antibiotic potentiating peptide #3.		176	4.60E-15
f5-14.aa	R69629	B. burgdorferi OspF operon.		106	3.50E-13
f5-14.aa	R89476	B. burgdorferi OspG lipoprotein.		157	6.20E-13
f5-14.aa	W26536	Trypanosoma cruzi antigen.		143	5.00E-11
f5-15.aa	R69629	B. burgdorferi OspF operon.		448	1.30E-68
f5-15.aa	R89476	B. burgdorferi OspG lipoprotein.		105	5.80E-24
f502.aa	R69852	Ethylene response (ETR) mutant protein etr1-3.		191	1.90E-35
f502.aa	R69849	Ethylene response (ETR) gene product.		191	2.70E-35
f502.aa	R69853	Ethylene response (ETR) mutant protein etr1-4.		191	2.70E-35
f502.aa	R69850	Ethylene response (ETR) mutant protein etr1-1.		191	3.60E-35
f502.aa	R69851	Ethylene response (ETR) mutant protein etr1-2.		191	3.60E-35
f502.aa	R74632	QETR ethylene response (ETR) protein from <i>Arabidopsis thaliana</i> .		190	5.20E-26
f502.aa	R74629	Tomato ethylene response (TER) protein.		171	6.50E-23
f502.aa	R74633	Nr (never ripe) tomato ethylene response (ETR) protein.		171	6.50E-23
f502.aa	R74630	Tomato TGETR1 ethylene response protein.		123	1.20E-19
f51-2.aa	W03626	Human thyrotropin GPR N-terminal sequence.		235	2.90E-23
f51-2.aa	R89476	B. burgdorferi OspG lipoprotein.		109	6.90E-23
f51-2.aa	W03627	Human follicle stimulating hormone GPR N-terminal sequence.		228	2.20E-22
f51-2.aa	W30763	Mannose-1-phosphate transferase protein MNM4.		203	1.00E-18
f51-2.aa	R70491	Leucocytozoan protozoa structural protein epitope.		191	7.50E-18
f51-2.aa	R97866	Chicken leucocytozoan immunogenic protein for use in vaccines.		183	4.80E-16
f51-2.aa	W21591	Antibiotic potentiating peptide #3.		159	6.20E-13
f51-2.aa	R68838	Plasmodium falciparum ABRA gene protein.		142	1.10E-12
f51-2.aa	R27530	Plasmodium falciparum blood and liver stage ABRA antigen.		142	2.80E-12
f51-2.aa	W31186	Human p160 polypeptide 160.2.		148	2.30E-11
f51-2.aa	W31185	Human p160 polypeptide 160.1.		148	2.40E-11
f517.aa	W24296	Staphylococcus aureus Gene #1 polypeptide sequence 2.		237	6.80E-30
f541.aa	R31013	P39-alpha.		1253	3.80E-229
f541.aa	R33280	P39-beta.		504	1.90E-117
f542.aa	R33280	P39-beta.		711	3.20E-96
f542.aa	R31013	P39-alpha.		101	7.90E-16

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f561.aa	R69631	B. burgdorferi T5 protein.	982	6.90E-131
f598.aa	W20289	H. pylori transporter protein, 24218968 aa.	264	9.90E-33
f598.aa	W20640	H. pylori transporter protein, 02ce110220orf8.	264	1.00E-30
f598.aa	W20101	H. pylori transporter protein 11132778 aa.	233	8.50E-27
f598.aa	W20861	H. pylori cell envelope transporter protein, 12ge10305orf16.	233	9.60E-27
f598.aa	W34202	Streptomyces efflux pump protein (frenolicin gene D product).	196	2.80E-21
f598.aa	R71091	C. jejuni PEB1A antigen from ORF3.	168	1.20E-17
f600.aa	W25527	Staphylococcus aureus Gene #20 polypeptide sequence 2.	209	3.40E-26
f600.aa	W34201	Streptomyces efflux pump protein (frenolicin gene C product).	169	6.50E-19
f600.aa	W20639	H. pylori transporter protein, 02ce110220orf7.	127	1.10E-14
f603.aa	W34200	Streptomyces efflux pump protein (frenolicin gene B product).	155	7.40E-32
f604.aa	R48035	Hyaluronic acid synthase of Streptococcus equisimilis.	110	2.30E-20
f606.aa	R48035	Hyaluronic acid synthase of Streptococcus equisimilis.	116	1.20E-25
f607.aa	R48035	Hyaluronic acid synthase of Streptococcus equisimilis.	141	1.50E-26
f631.aa	W15192	Staphylococcus aureus cell surface protein.	160	7.30E-29
f664.aa	W20105	H. pylori flagella-associated protein, 1171928 aa.	202	3.20E-46
f664.aa	W20688	H. pylori flagella-associated protein 04ge11713orf5.	202	2.60E-45
f664.aa	R97245	Virulence gene cluster polypeptide product.	158	3.90E-13
f704.aa	R60153	Nematode-inducible transmembrane pore protein.	104	2.50E-18
f704.aa	R33913	Sequence encoded by TobRB7-5A which encodes a membrane channel	104	2.50E-18
f704.aa	R77082	Tobacco root specific promoter RB7 from clone lambda5A (TobRB7-5A).	104	2.50E-18
f742.aa	W46499	Amino acid sequence of the spindly (SPY) protein of Arabidopsis.	101	2.50E-14
f752.aa	W20733	H. pylori cell envelope protein, 06cp11722orf15.	141	3.00E-37
f752.aa	W20358	H. pylori cell envelope protein 26366312 aa.	110	4.20E-18
f814.aa	W20753	H. pylori transporter protein, 06gp112020orf7.	178	7.90E-35
f814.aa	W20420	H. pylori cell envelope transporter protein 33399142.aa.	160	2.30E-21
f843.aa	R14319	Human T-cell immunosuppressive factor.	167	1.20E-19
f860.aa	W21894	Asparaginyl-tRNA synthetase from <i>Staphylococcus aureus</i> .	245	2.30E-38
f860.aa	W33903	Streptococcus pneumoniae asparaginyl tRNA synthetase.	177	1.10E-22
f867.aa	W34261	An alpha subunit of a thermostable ATPase.	592	1.30E-161
f867.aa	R10098	Alpha subunit of ATP-synthase.	741	4.90E-144

TABLE 2. Closest matching sequences between the polypeptides of the present invention and sequences in GenBank and Derwent databases.

f867.aa	R31522	Carrot reverse transcriptase.	311	4.60E-130
f867.aa	R10099	Beta subunit of ATP-synthase.	121	7.90E-14
f867.aa	W34262	A beta subunit of a thermostable ATPase.	116	1.00E-12
f868.aa	W34262	A beta subunit of a thermostable ATPase.	151	6.10E-109
f868.aa	R10099	Beta subunit of ATP-synthase.	172	1.90E-106
f868.aa	W34261	An alpha subunit of a thermostable ATPase.	117	3.10E-19
f868.aa	R10098	Alpha subunit of ATP-synthase.	113	2.00E-18
f868.aa	R31522	Carrot reverse transcriptase.	101	7.10E-15
f874.aa	R10591	L-lactic acid dehydrogenase.	538	7.20E-109
f874.aa	R08355	Recombinant thermophilic NAD-dependant dehydrogenase.	455	9.80E-99
f874.aa	R09295	Recombinant thermophilic NAD-dependant dehydrogenase.	455	9.80E-99
f874.aa	R15736	L-lactic acid dehydrogenase.	426	1.60E-85
f874.aa	P91948	Pig H4 isoenzyme.	393	2.00E-82
f874.aa	W33108	Chicken lactic acid dehydrogenase type B subunit.	390	2.20E-80
f874.aa	W33107	Chicken lactic acid dehydrogenase type B subunit.	385	1.10E-79
f874.aa	P80891	Testis-specific lactate dehydrogenase subunit LDH-C4.	339	5.50E-74
f874.aa	R94013	Heat resistant maleate dehydrogenase.	255	1.30E-55
f874.aa	R11119	Recombinant L-2-hydroxyisocaproic acid dehydrogenase.	224	7.90E-49
f874.aa	R62605	P. falciparum lactate dehydrogenase.	255	2.00E-44
f874.aa	W11476	Emmeria lactate dehydrogenase.	203	1.10E-25
f943.aa	P91223	Coenzyme-independent L-sorbose dehydrogenase from Gluconobacter	175	4.30E-16

TABLE 3. Conservative Amino Acid Substitutions.

Aromatic	Phenylalanine Tryptophan Tyrosine
Hydrophobic	Leucine Isoleucine Valine
Polar	Glutamine Asparagine
Basic	Arginine Lysine Histidine
Acidic	Aspartic Acid Glutamic Acid
Small	Alanine Serine Threonine Methionine Glycine

TABLE 4. Residues Comprising Epito-Bearing Fragments

Query	Residues Comprising Epito-Bearing Fragments
f101.aa	from about Lys-62 to about Gly-64, from about Ser-111 to about Asp-113, from about Arg-136 to about Arg-139, from about Pro-189 to about Asn-193.
f11.aa	from about Pro-38 to about Lys-40, from about Glu-92 to about Lys-96.
f12.aa	from about Pro-288 to about Asp-290, from about Asn-336 to about Gly-338, from about Tyr-410 to about Gly-413, from about Asp-418 to about Arg-420, from about Pro-552 to about Val-555, from
	about Gln-643 to about Asp-645, from about Gln-1061 to about Arg-1063, from about Asn-1130 to about Lys-1132.
f129.aa	from about Glu-76 to about Arg-81, from about Lys-144 to about Asn-146.
f147.aa	from about Gln-94 to about Thr-96.
f152.aa	from about Gly-35 to about Gly-37, from about Gln-321 to about Gly-323.
f154.aa	from about Asn-39 to about Lys-41, from about Ser-74 to about Lys-77, from about Ser-213 to about Gly-215, from about Ser-303 to about Asp-306, from about Asp-422 to about Asn-424.
f157.aa	from about Lys-21 to about Asp-24, from about Ser-45 to about Tyr-47.
f17.aa	from about Arg-17 to about Asn-20, from about Thr-94 to about Gly-96.
f186.aa	from about Lys-305 to about Tyr-308.
f196.aa	from about Lys-121 to about Asn-123, from about Pro-278 to about Lys-282, from about Glu-576 to about Tyr-578.
f899.aa	from about Asn-174 to about Asp-177.
f925.aa	from about Lys-201 to about Asp-204, from about Phe-291 to about Lys-294.
f929.aa	from about Pro-139 to about Asn-141, from about Arg-211 to about Glu-214, from about Thr-370 to about Asn-375.
f933.aa	from about Ser-139 to about Lys-143.
f940.aa	from about Gly-143 to about Asn-148.
f943.aa	from about Asp-58 to about Asp-60, from about Lys-157 to about Asn-159, from about Asp-217 to about Asp-221, from about Lys-250 to about Asn-254, from about Pro-262 to about Asn-264, from about Gly-305 to about Trp-307.
f952.aa	from about Ser-52 to about Ser-54.
f4.aa	from about Arg-64 to about Arg-67.
f43.aa	from about Ser-84 to about Gln-87, from about Asp-231 to about Tyr-233, from about Arg-296 to about Asp-300.
f50.aa	from about Glu-136 to about Gly-138, from about Asp-153 to about Lys-155, from about Asp-289 to about Asp-291, from about Glu-458 to about Asn-461.
f65.aa	from about Glu-120 to about Asp-122, from about Pro-204 to about Tyr-206.
f8.aa	from about Pro-263 to about Arg-265, from about Asp-274 to about Lys-278.
f82.aa	from about Tyr-66 to about Gly-68, from about Ser-116 to about Lys-119, from about Asp-121 to about Gly-123, from about Pro-128 to about Gly-131.

TABLE 4. Residues Comprising Epito-Bearing Fragments

f86.aa	from about Asn-179 to about Asn-181, from about Lys-192 to about Asn-194, from about Lys-270 to about Asn-272, from about Lys-279 to about Lys-282, from about Asp-331 to about Asn-333.
f477.aa	from about Pro-250 to about Lys-253.
f488.aa	from about Lys-76 to about Lys-79, from about Asn-486 to about Asp-489, from about Lys-508 to about Gly-510, from about Asn-559 to about Gly-562.
f494.aa	from about Lys-76 to about Asn-78.
f516.aa	from about Lys-32 to about Asp-34.
f523.aa	from about Pro-202 to about Asn-206, from about Lys-255 to about Tyr-258.
f526.aa	from about Asn-85 to about Lys-88, from about Asp-136 to about Gly-138.
f577.aa	from about Cys-18 to about Lys-22, from about Asn-297 to about Gln-300.
f584.aa	from about Pro-131 to about Lys-133, from about Pro-200 to about Ser-202.
f596.aa	from about Arg-42 to about Asp-44, from about Asp-117 to about Tyr-119, from about Pro-205 to about Asp-207.
f600.aa	from about Pro-143 to about Asp-145.
f603.aa	from about Phe-35 to about Ser-37.
f607.aa	from about Gln-67 to about Lys-70, from about Asp-273 to about Tyr-275, from about Asp-333 to about Gly-338, from about Pro-359 to about Lys-362, from about Arg-409 to about Gly-411.
f611.aa	from about Arg-133 to about Gly-135.
f631.aa	from about Pro-132 to about Asn-136, from about Asn-159 to about Tyr-161, from about Pro-216 to about Asp-218, from about Pro-220 to about Lys-223.
f688.aa	from about Lys-266 to about Asp-268, from about Lys-271 to about Asn-273, from about Lys-315 to about Lys-318.
f704.aa	from about Lys-250 to about Lys-253.
f707.aa	from about Lys-131 to about Asp-134, from about Asp-246 to about Asn-249.
f709.aa	from about Tyr-39 to about Gly-42, from about Lys-148 to about Gly-150, from about Arg-269 to about Gly-272, from about Ser-466 to about Tyr-468, from about Asn-489 to about Asn-491, from about Lys-575 to about Asp-578, from about Pro-642 to about Lys-644.
f197.aa	from about Pro-217 to about Asp-219, from about Glu-675 to about Asp-678, from about Pro-687 to about Asn-689, from about Glu-694 to about Gln-696.
f200.aa	from about Arg-174 to about Phe-179.
f208.aa	from about Arg-326 to about Ser-328.
f210.aa	from about Pro-191 to about Ile-194.
f221.aa	from about Asn-133 to about Asn-135.
f253.aa	from about Arg-191 to about Gly-194.
f269.aa	from about Ser-271 to about Thr-273, from about Asp-284 to about Gly-286.
f29.aa	from about Pro-159 to about Ser-161.
f290.aa	from about Pro-240 to about Gly-244.
f291.aa	from about Gln-267 to about Lys-269.

TABLE 4. Residues Comprising Epito-Bearing Fragments

f296.aa	from about Glu-98 to about Lys-101.
f3.aa	from about Asn-241 to about Lys-245.
f30.aa	from about Asn-156 to about Tyr-159, from about Asn-178 to about Lys-180.
f939.aa	from about Ser-245 to about Asn-249.
f739.aa	from about Asn-80 to about Tyr-82, from about Lys-208 to about Ser-210.
f742.aa	from about Ser-141 to about Asp-145, from about Asn-222 to about Gln-225, from about Asp-243 to about Tyr-247, from about Asn-249 to about Asn-251.
f743.aa	from about Arg-111 to about Gly-114, from about Pro-131 to about Asp-134.
f790.aa	from about Thr-40 to about Asn-42, from about Ser-53 to about Ser-55, from about Lys-215 to about Asp-218, from about Asn-274 to about Gly-277.
f792.aa	from about Val-82 to about Ser-84, from about Ser-102 to about Asn-104, from about Gln-127 to about Tyr-130, from about Lys-309 to about Asn-314, from about Lys-375 to about Thr-377, from about Pro-511 to about His-513, from about Thr-515 to about Asp-517.
f797.aa	from about Pro-119 to about Gly-122, from about Lys-166 to about Asn-169.
f799.aa	from about Asn-31 to about Asn-34, from about Gln-44 to about Asn-47, from about Pro-123 to about Gly-125.
f814.aa	from about Ser-120 to about Ser-122, from about Arg-636 to about Asn-638, from about Cys-967 to about Ser-969.
f820.aa	from about Thr-563 to about Tyr-565.
f850.aa	from about Tyr-159 to about Tyr-164, from about Gln-375 to about Asp-379.
f853.aa	from about Thr-180 to about Lys-184, from about Arg-231 to about Asp-233, from about Asn-252 to about Gly-254.
f859.aa	from about Lys-46 to about Ser-52, from about Pro-88 to about Asn-91; from about Asn-117 to about Asp-120.
f861.aa	from about Asp-38 to about Lys-40, from about Lys-219 to about Asn-225.
f368.aa	from about Gln-228 to about Asn-231.
f371.aa	from about Tyr-109 to about Asn-111, from about Asn-162 to about Gln-164.
f502.aa	from about Asn-118 to about Lys-122, from about Ser-269 to about Gly-271, from about Lys-370 to about Asp-373, from about Asn-509 to about Lys-511, from about Lys-705 to about Arg-707, from about Thr-912 to about Gly-914, from about Pro-1213 to about Asp-1216, from about Asn-1491 to about Arg-1493.
f527.aa	from about Cys-20 to about Gln-22, from about Asn-38 to about Asn-40, from about Phe-112 to about Asp-114, from about Lys-160 to about Asn-162, from about Ser-199 to about Asp-201, from about
	Gln-258 to about Gly-261, from about Arg-282 to about Asn-284, from about Ser-297 to about Asp-299.
f541.aa	from about Ser-68 to about Asn-71.
f604.aa	from about Lys-77 to about Gly-79, from about Lys-201 to about Asn-203, from about Asp-252 to about Asp-254, from about Tyr-

TABLE 4. Residues Comprising Epito-Bearing Fragments

	347 to about Gly-350, from about Asp-514 to about Trp-516.
f736.aa	from about Lys-20 to about Asn-24, from about Arg-147 to about Ser-153, from about Ser-231 to about Lys-233.
f752.aa	from about Thr-119 to about Lys-122, from about Pro-420 to about Gly-422.
f798.aa	from about Asp-33 to about Thr-36, from about Lys-180 to about His-183.
f635.aa	from about Pro-100 to about Asn-102, from about Asp-145 to about Phe-147.
f32.aa	from about Lys-18 to about Asn-20.
f320.aa	from about Asn-193 to about Leu-195, from about Gln-248 to about Lys-250.
f352.aa	from about Ser-46 to about Asn-49.
f301.aa	from about Lys-178 to about Lys-180, from about Ser-401 to about Tyr-404.
f373.aa	from about Gly-88 to about Lys-90, from about Asn-539 to about Lys-542, from about Glu-654 to about Ser-657.
f384.aa	from about Pro-250 to about Asn-252, from about Asp-266 to about Lys-268.
f446.aa	from about Asp-20 to about Ser-26, from about Asn-146 to about Lys-149.
f542.aa	from about Arg-86 to about Gly-88, from about Arg-163 to about Asn-165.
f93.aa	from about Asn-152 to about Asp-155.
f105.aa	from about Asp-48 to about Phe-50.
f150.aa	from about Thr-214 to about Asp-218, from about Asp-256 to about Asp-259.
f219.aa	from about Asn-77 to about Asn-81, from about Asp-111 to about Asn-115.
f229.aa	from about Gln-61 to about Asn-63.
f32.aa	from about Lys-18 to about Asn-20.
f186.aa	from about Lys-305 to about Tyr-308.
f216.aa	from about Ser-105 to about Asn-107.
f328.aa	from about Asn-105 to about Asp-107.
f352.aa	from about Ser-46 to about Asn-49.
f867.aa	from about Thr-3 to about Gly-5, from about Lys-156 to about Ser-159.
f868.aa	from about Arg-94 to about Gly-96, from about Pro-257 to about Gly-261, from about Pro-295 to about Asp-297, from about Arg-340 to about Asp-342.
f872.aa	from about Ser-19 to about Lys-23, from about Thr-139 to about Asp-142, from about Ser-282 to about Tyr-286, from about Ser-311 to about Ser-313.
f886.aa	from about Thr-83 to about Asp-85, from about Asp-106 to about Lys-108, from about Lys-143 to about Gly-147, from about Asp-186 to about Asn-191.
f888.aa	from about Asn-65 to about Asp-67.
f893.aa	from about Asn-203 to about Asn-207, from about Thr-446 to about Asn-450.
f605.aa	from about Arg-31 to about Asp-33.
f606.aa	from about Asn-68 to about Gly-71, from about Asn-136 to about

TABLE 4. Residues Comprising Epito-Bearing Fragments

	Lys-139, from about Asn-223 to about Tyr-226, from about Ser-276 to about Tyr-279, from about Pro-362 to about Asn-365, from about Arg-503 to about Trp-507.
f679.aa	from about Lys-154 to about Asp-156, from about Lys-224 to about Arg-226, from about Asn-260 to about Asp-264, from about Glu-363 to about Lys-366, from about Asp-387 to about Gly-389, from about Tyr-441 to about Lys-443, from about Arg-501 to about Tyr-504.
f11-12.aa	from about Pro-91 to about Asn-93, from about Pro-181 to about Asp-186, from about Lys-244 to about Ser-248.
f11-4.aa	from about Asn-160 to about Lys-163.
f14-8.aa	from about Pro-92 to about Gln-95, from about Lys-123 to about Thr-125, from about Lys-215 to about Asp-219.
f17-6.aa	from about Pro-36 to about Glu-38.
f19-2.aa	from about Ser-104 to about Ser-106, from about Gln-230 to about Asn-232.
f19-4.aa	from about Val-79 to about Thr-82, from about Pro-195 to about Gly-201.
f19-6.aa	from about Asp-24 to about Lys-30, from about Pro-36 to about Glu-38.
f21-4.aa	from about Cys-24 to about Asn-26.
f28-2.aa	from about Ser-77 to about Lys-80, from about Tyr-274 to about Asn-277.
f28-3.aa	from about Glu-53 to about Arg-57, from about Gln-82 to about Asn-85, from about Gln-157 to about Asn-159.
f31-2.aa	from about Arg-95 to about Arg-97, from about Asn-297 to about Asn-299.
f4-15.aa	from about Pro-182 to about Asp-184, from about Lys-220 to about Asp-222.
f4-50.aa	from about Thr-109 to about Asn-111.
f42-1.aa	from about Asn-55 to about Asn-57, from about Arg-81 to about Ser-84, from about Asp-94 to about Asn-97.
f45-2.aa	from about Asn-83 to about Gly-86.
f47-2.aa	from about Ser-29 to about Asp-33, from about Asn-94 to about Lys-99, from about Pro-152 to about Lys-157.
f49-2.aa	from about Asn-452 to about Gly-454.
f5-14.aa	from about Glu-102 to about Asp-106, from about Thr-272 to about Asn-275, from about Glu-313 to about Asn-315, from about Ser-370 to about Ser-372.
f5-15.aa	from about Lys-170 to about Gly-173, from about Asn-194 to about Gly-196.
f51-2.aa	from about Asp-302 to about Lys-304.
f6-21.aa	from about Glu-38 to about Asn-42, from about Arg-84 to about Gly-87.
f6-27.aa	from about Asp-67 to about Asn-69, from about Arg-85 to about Asn-89, from about Lys-168 to about Gly-171, from about Lys-179 to about Asn-181, from about Ser-380 to about His-382.
f6-5.aa	from about Ser-67 to about Asn-71.
f7-30.aa	from about Pro-94 to about Asp-96, from about Lys-144 to about Arg-147.
f76-1.aa	from about Asn-30 to about Lys-35, from about Lys-113 to about

TABLE 4. Residues Comprising Epito-Bearing Fragments

	Gly-116, from about Glu-119 to about Lys-121.
f8-10.aa	from about Pro-25 to about Lys-32, from about Ser-168 to about Thr-172.
f01a.aa_bb001	from about Pro-123 to about Asp-125, from about Ser-179 to about Asp-181, from about Lys-255 to about Gly-259.
_bb0011	from about Ala8 about Arg 17, from about Tyr31 to about Gly40, from about Ser65 to about Lys78, from about Val93 to about Asp102, from about Ser120 to about Ile129, from about Pro156 to about Glu170, from about Lys187 to about Asn 196, from about His205 to about Lys214, from about Gly226 to about Glu235, from about Gln253 to about Asn266, from about Glu283 to about Glu293, from about Leu311 to about Ile320, from about Arg326 to about Gly335, from about Pro340 to about Ala349
f02a.aa_bb002	from about Tyr-169 to about Asn-171, from about Tyr-242 to about Asn-245, from about Lys-264 to about Asp-267.
_bb9	from about Met7 to about Lys16, from about Lys47 to about Ser57, from about Asn80 to about Ser89, from about Gly103 to about Glu113, from about Lys125 to about Pro133, from about Lys138 to about Ala147
f03a.aa_bb006	from about Asp-54 to about Thr-57, from about Lys-201 to about His-204.
_bb014	from about Pro23 to about Gln31, from about Ser37 to about Asp45, from about Leu76 to about Asn84, from about Leu76 to about Val84, from about Ser89 to about Asn97, from about Ser105 to about Lys113, from about Asn120 to about Met128, from about Asn159 to about Gly 167, from about Lys173 to about Bal181
_bb023	from about Asp17 to about Gly27, from about Arg40 to about Asp48, from about Val64 to about Asp72, from about Glu105 to about Thr113, from about Ser141 to about Gly150, from about Asp155 to about Ile163, from about Asn184 to about Lys198, from about Ile219 to about Pro227, from about Ser230 to about Phe238, from about Ser241 to about Asn250, from about Asp270 to about Val278, from about Ser285 to about Leu293, from about Glyu307 to about Ser315, from about Lys327 to about Asn335
f08a.aa_bb024	from about Asn-30 to about Asp-33, from about Ser-116 to about Asn-118, from about Asn-154 to about Gly-156.
f09a.aa_bb025	from about Asn-30 to about Ser-35, from about Thr-145 to about Asn-148.

Applicant's or agent's file reference number	PB3072	International application	Unassigned
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INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

<p>A. The indications made below relate to the microorganism referred to in the description on page <u>8</u>, line <u>8</u></p>			
<p>B. IDENTIFICATION OF DEPOSIT</p>		Further deposits are identified on an additional sheet <input type="checkbox"/>	
Name of depositary institution American Type Culture Collection			
Address of depositary institution (<i>including postal code and country</i>) 10801 University Boulevard Manassas, Virginia 20110-2209 United States of America			
Date of deposit August 8, 1998		Accession Number 202012	
<p>C. ADDITIONAL INDICATIONS (<i>leave blank if not applicable</i>) This information is continued on an additional sheet <input type="checkbox"/></p>			
<p>D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (<i>if the indications are not for all designated States</i>)</p>			
<p>E. SEPARATE FURNISHING OF INDICATIONS (<i>leave blank if not applicable</i>)</p> <p>The indications listed below will be submitted to the International Bureau later (<i>specify the general nature of the indications, e.g., "Accession Number of Deposit"</i>)</p>			
<p>For receiving Office use only</p> <p><input type="checkbox"/> This sheet was received with the international application</p>		<p>For International Bureau use only</p> <p><input type="checkbox"/> This sheet was received by the International Bureau on:</p>	
Authorized officer <div style="height: 40px; border: 1px solid black; margin-top: 5px;"></div>		Authorized officer <div style="height: 40px; border: 1px solid black; margin-top: 5px;"></div>	